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SARKIS BOSHNAKIAN
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THE GENETICS OF SQUAREHEADEDNESS AND OF DENSITY IN WHEAT, AND THE RELATION OF THESE TO OTHER CHARACTERS

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THE GENETICS OF SQUAREHEADEDNESS AND OF \\ DENSITY IN WHEAT, AND THE RELATION OF THESE TO OTHER CHARACTERS \({ }^{1}\)
}

\section*{Sarkis Boshnakian \({ }^{2}\)}

The mendelian inheritance of each of the more striking characters in wheat, such as beardedness, color, felting, density, and so forth, has been determined qualitatively by various workers. Much remains to be done, however, if the genetics of these characters is to be analyzed from a quantitative point of view, as there are numerous lesser inherited variations within their simple gross segregations.

Practically all of these characters show certain degrees and types of interrelations with other characters. Some of them show complete or partial linkage. Others, when analyzed quantitatively, appear to have been affected by one or another character but not necessarily linked with it, so that if one of these characters appears in an intense or a dilute form the others vary also in the same direction and more or less in the same degree.

Besides the above-mentioned type of interrelation, in which the appearance and the intensity of appearance of a group of characters are due to the presence or the absence of the same factor or factors, there is another type embracing a set of characters that appear as an indirect effect of the presence of another character. In a sense these characters are acquired, but they seem to be inherited simply because the causal character is inherited; and whenever the latter is present it gradually causes the modification of the former characters during the lifetime of the individual.

The subject of the inheritance of types of internode length presented in this paper has been treated from the following three viewpoints: (I) the analyses of minor segregations within gross \((3: I, I: 2: 1\), or the like) segregations; (2) the determination of the interrelations of varied

\footnotetext{
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\({ }^{2}\) In cooperation with the Office of Cereal Investigations, United States Department of Agriculture.
}
characters; and (3) the determination of characters that were found to be the resultant of other characters.

Since the characters studied were confined to those that were not distinctly contrasting in the usual mendelian sense but represented different gradations on a scale between two extreme quantitative characters, it was not possible to classify them into genetic classes or to express the results always in terms of ratios. The analyses were made according to such biometrical methods as seemed best suited to bring out the directions and tendencies of the variations. Factorial explanations, however, have been given wherever the facts obtained warranted the formulation of such hypotheses.

The material on which observations were made consisted, exclusive of interspecific crosses, of more than sixty \(\mathrm{F}_{2}\) progenies, fourteen of which were carried through the \(\mathrm{F}_{3}\). To avoid duplications of similar results it is not considered necessary to present here the results of all the crosses, but sufficient data are given to serve as illustrations and to show the general trend of the various modes of inheritance.

For many valuable suggestions and criticisms during the progress of this work the writer fully acknowledges his indebtedness to Professor H. H. Love, of the Department of Plant Breeding at Cornell University, under whose direction the studies were made. Most of the \(\mathrm{F}_{2}\) and \(\mathrm{F}_{3}\) progenies studied were furnished by Dr. C. E. Leighty, of the United States Department of Agriculture. The writer wishes to express his appreciation of Dr. Leighty's generosity in supplying these and also carefully numbered hybrid progenies, which made possible the tracing of each back to the \(F_{1}\) and parental material, all of which had been saved. Thanks are due also to the Office of Cereal Investigations, United States Department of Agriculture, through whose courtesy the writer was enabled to enjoy the field and laboratory facilities at the Arlington Experimental Grounds during the summers of 1916 and 1917.

\section*{PHYSIOLOGICAL CONDITIONS AFFECTING RACHIS INTERNODE LENGTH}
like many quantitative characters, density and squareheadedness are affected to a greater or less extent by a number of environmental conditions which tend to change the normal course of development of the plant, thereby suppressing or accelerating the growth of certain of its
parts. A brief discussion of the effect of environmental factors on the production of these internode characters will serve to show to what extent nongenetic variations may take place. In the first part of this paper the main object is to explain the phenomena of density and squareheadedness on a purely mechanical basis.

\section*{DETERMINATION OF DENSITY AND OF SQUAREHEADEDNESS}

The terms density and squarcheadedness are used in this paper to designate two different phenomena of rachis internode length. The differences between these two characters are discussed elsewhere (Bushnakian, 1917) \({ }^{3}\), but they may be briefly redescribed here.

Density is expressed in terms of average rachis internode length, which is found by dividing the length of the rachis by the number of rachis internodes. The average internode length, or density, of a heard 90 millimeters long with 20 rachis internodes, is thus \(90 \div 20\), or 4.5 millimeters. The average internode length in wheat varies from about I. 3 to about \(S\) millimeters. In z'ulgare wheat it usually does not extend beyond 5 millimeters. Density is comparative.

The average internode length usually varies in dense wheats from i. 2 to 2.5 millimeters, and in lax forms from 3 to \(5+\) millimeters. There are intermediate gradations, but, in a general way, by dense or lex forms are meant, respectively, forms varying in density between the two ranges specified. Although the word club refers to a capitate type of head, following the present tisage of this term it is here applied to demse wheats of the sutizan group (that is, Triticum compactum) whether capitate or not.

Squareheadedness, on the other hand, refers to the ratio between the average internode length of the central third of the rachis and that of the temmal third. The density of the middle and nrper thirds of the rachis is equal to the ratio of the number of internodes in these two sections of the rachis. The coefficient of squareheadedness is found by dividing the number of rachis internodes in the tominal third of the rachis by the number of internodes in the middle third. 'Thus, if the terminal third has 9.5 internodes and the central third has 5.6 internodes, the degree of squareheadedness is equal to \(9.5 \div 5.6\), or 1.69 . The forms
\({ }^{3}\) Dates in parenthesis refer to Literature Cited. page 881.
that are usually called squareheads have a coefficient of about I. 33 or more; and in this paper, by vulgare or non-squarchead forms are meant types with coefficients of less than I.33, and by squareheads, forms having coefficients of more than I.33. This division is entirely arbitrary and is made for convenience.

It is seen, then, that squarcheadedness has no reference to density. Squareheads may be dense or lax. This divides the wheat types into the following font classes with respect to their rachis internodes: nonsquarehead, lax (Triticum vulgure, Plate LNVII, upper, I2) ; squarehead, lax (Tr. capitatum, Ilate LNVII, upper, II ) ; non-squarehead, club (Tr. compactum, Plate LNVII, lower, I3) ; squarehead, club (Tr. com-pacto-capitatum Plate LNVII, upper, 9). These varietal names apply to the wheats of the sativum group only.

According to these classifications, the semi-dense forms having a squareheadedness of less than I. 33 are named semi-dense vulgare-not squarelieads, a name too often applied for such forms, and perhaps with some justification as such semi-dense forms when well developed may appear square in cross section.

Since the discussions in this paper center solely tupon squareheadedness and density, it is necessary for the reader to bear in mind the sense in which these two terms are used.
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DFVELOPMENT OF THE WHEAT PLANT WITH KEFERENCE TO SQUARETEADEDNESS

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The head of the wheat plant is found in an embryonic stage when the plant starts a new growth after a short or a long period of rest. When the head is about io millimeters long it is covered with concentric rings of sheaths, each sheath being attached to the culm at its respective node. The different parts of the plant do not all grow at the same rate.

When the spike is about 15 millimeters long, the enveloping sheaths and blades are fairly well developed, but the culn internodes are only a few millimeters in length, the terminal ones being the shortest. After the sheaths have gone through their chief period of growth, the development of the internodes is accelerated. During this period the head also begins to develop. From this time on, the increase in the height of the


VARIOUS FORMS OF GROWTH
Upper: 1, Aegilops ovata: 2, Fi (Aegilops x Silver Club); 3, Silver Club. 4, White Spelt; 5, Dale Gloria; 6 to 12, F'z types (White Spelt x Dale Gloria, series 13255a) - 6, homozygous lax spelt; 7 , heterozygous deasp spelt. 8 . homozygous dense spels; 9, homozygous deuse sativum (club): 10, heterozygous dense sataum; 11, homozygous lax sativum, squarehead; 12, homozygous lax sativum, vulgare
Lower: Dense forms of different species: 1, durum; 2 and 3, capitate und dene barley heads (lateral florets of 2 removed); 4, capitate sativum; 5, dense spelt; 0 , dense polonetum; 7 and 8 , elub wheats; 9, furgidum; 10, club; 11 and 12, capitate dicoccums; 13, club pyramidal; 14, capitate rpelt; 15, dense polorucum; 16, club


VARIOUS FORMS OF GROWTH
Upper: 1 to 5, 6 to 10 , Heads of two hybrid plants showing lack of harmony of growth characters. 11 and 12, Heades (of spelts subjected to longitudinal pressure: zigzagging of internodes produced instead of squareheadedness
Lower: 1 to 4, 5 to 8. Heads of two plants showing variations in density on mame plant; plants grown in greeshouse; long spikes headed out about four weeks earlier than dense epikes
plant is due primarily to the increasing length of each culm internode. During the first period of the development of the culm, the basal internodes, except a few near the ground, begin to develop, and successive internodes undergo their chief period of growth as the growth of the internode below is beginning to decline. The terminal section of the culm, which carries the spike, during its development has to push the spike up all along the length of the temmal heath, which enselog the head in such a way that occasionally, and especially among plants of hybrid origin, the culm cannot exert sufficient pressure to unfold the sheath. In such cases the spike fails to head out; or, if it finally does head wot, the head apmears in an almormal condition and the tip yitielets very often remain undeveloped (Plate LdVIII, upper, I to IO).

The factors that seem to produce a strain on the terminal culm internode during its growth are the following: the rapid growth of the culm; the spread, width, shape, and texture of the spikelets; the stiffness of the enveloping sheath and its resistance to unfolding.

It seems that as the rate of growth of the terminal culm increases, the anonement of the spike through the sheath should enconnter atseater degree of resistance in an opposite direction.

The spread of the spikelets is probably one of the most important factors. The glumes of the spikelets are pointed upward and outward. This in itself tends to increase resistance. If the contact of the glumes with the sheath increases their spread, the resistance will ingrease many fold. In species such as the spelt or the emmer, in which the spikes are very narrow and the spikelets are very close to the rachis, the resistance is decidedly decreased because the spikelets themselves assume a wedge shape, the glumes being drawn together tightly; and also because the spikelets, lying flat against the rachis, are not likely to spread out.

The third factor, which is not so important as the other two, is the texture of the sheath and its habit of development. The sheath that normally unfolds at the proper time, or is easily unfolded by the movement of the spike, sometimes fails to open completely or opens under difficulty. Sheathe uf thin type produce a com-ibleralle kongitudinal pressure on the culm.

In wrder to understand the mechance of the production of squareheadednes. it is necessary to know the effect that in produced ly pressure aloner the culm axis. The presence of longitudinal pressure is evident from the undulations of the culms often observed in square or dense forms : Plate LXIII, lower. 9 and Ito. 1 The pert of the spike that most react- to the effect of presotre is the terminal part, because it is directly in contact with the sheath. The pressure producen a compressing effect. and this in turn checks the development of the terminal part of the head. especially the development of the rachic internoles and produces the effect known as squareheadedness.

This character of squareheadednes is mainly evidenced by the gradua: Shortening of the terminal rachis internodes , Plate LXITI, lower. f. If, and 12. But there are also other characters which accompany this shortenines of the internoles and which are the direct or indirect results of the same catuse. One of the most promment of the ere is the so-called chubbing. or capitate form, probluced be the spreading of the spikelets away foms the rachis in those regons of the head where the internodes are wort Becathe of the presure exoled, the normal elongation of the internodeis inhibited but the spikelets in most cases continue to develop. Since the-pare between the ppikelet is mot sutticient. they are forced mechanically to pread out to make more rosm for derelonment. 'Thn proses is on the principle of the iswacele traperobld, in which, the lave being constant, the distance between the siden increase it the latter take a position toward a right angle with the base. In the plant the rachis internole in repreented by the base, and the axes of the spikelet by the sides, of the trapezoid.

In squareheads the spikelets of only the upper part of the spike (except the terminal two or three spikelets) thus diverge. In most comfactum forms all spikelet diverge a- a renth of the -horthe- wifl the intertomber This is seen on comparing the divergence of prikelets of dence heats 5 and 0 . in I'late I. STII 1 uptrer , with that of las head= 2 . 4, and 6. In wheat, as well as in barley, the opposite condition exists
 near the upper part, but near the basal region. In such cases the ear, in-

St sad ut being capitate in form, ashumes a pramidal or conical form, az shown he heads I3. Plate L.

Another phenomenon of sytureheadedness is the drawing of the terminal spikelets toward one side so that when the head is viewed ahner one of the directions of the plane of symmetry, which separates the spitelets of one side from those of the other, the rachis appears exposed (Plate LXVII, upper, II). Viewed from the opposite side the rachis is covered by glumes and awns which are outdrawn and gathered in that dircetion. With the receding of the glames the part of the side where the :achis is exposed appears that ( Plate LXIII, upper, II), and to a lesan not vewing the head from the opposite side also it gives the imp:ession that the spike is square in cross section. This impression, which has been left on the popular mind, has given to this form the name squarehead.

The character of squareheadedness is not confined to the wheat known by this name but may appear also in the dense forms known as club wheats , Triticum compactum). The total shortening of rachis internodes in these forms is primarily due to the presence of a genetic factor which produces aloo a general shortening of many parts of the plant. But in nurs forms a certain degree ni sytareheadednes may he found. This may ie inherent-that is, transmissible-s it may have been produced mechanically: When the spikclete spread out as a revult of the hortnesani the internodes, as decribed alowe, the increased width of the heat and the projections of the tipe of the glume are likely to offer consiterabie resistance, thus producing squareheadednese in the manner already explained.

\section*{EFFECTS OF CROSSING ON SQUAREHEADEDNESS}

Syuareheadedness, and shortening of all the rachis internodes are two different phenomena. As shown in the preceding discussion, supareheadedness is a postnatal character, as it were, being dependent on the combined effect of certain vegetative growth characters. Density of the compactum wheats, on the other hand, is predetermined and is due to the presence of one or more genetic factor which catuse dwarfing if a numise: of plant parts, including incilentally the shortening of all rachis internodes.

Sypareheadedness is dependent no a eertain balance of the rate of
growth of the parts concerned. An unfavorable balance produced through hybridization may result in certain hereditary anomalies. A few such forms are shown in Plate LAVIII (upper). Heads I to 5 were produced on a single \(F_{2}\) plant derived from a cross between a durum and a common wheat. In these cases the curling of the awns all along the length of the heads shows that the latter were partly prevented from men ing up the theath- lay the hemus of the theath. The illu-trations show also the rudimentary condition of the terminal 5 to 7 spikelets, which represent the region whose growth was checked altogether by being subjected to pressure.

Heads 6 to 10 in the same plate represent another condition of lack of harmony of growth between different parts of the plant. The spiral form of the awns of head 7 shows that this head was forced to make a corkscrew movement while making its way up the sheath. Heads 8 and Io show the failure of the sheath to open at the proper time. Heads 6 and 9 represent heads that were finally released.

Heads II and 12 represent a single spelt plant whose sheaths were evidently wrapled tue tightly arnumb the head. The pres-ure which the tight sheath exerted on the head by the growth of the culm produced a zigzagging of the rachis. The internodes of the spelts, being comparatively stiff, are not so likely to remain short as a result of pressure.

These two spelt heads are interesting because they show the relative tendency of the different internodes to be affected by pressure. The basal internodes are thick and are very slightly affected by the induced zigzagging effect. Each successive internode is weaker than the one below, and more and more likely to show the effect of pressure. The conditions to which these heads were subjected are identical with those to which squareheads of satizum or other soft-glumed species are subjected, but the effect is somewhat different because of the differences of texture and ear form of the spelt as compared with those of some other forms.

These cases show that there are a number of growth characters to which the production of squareheadedness is due, and that the factors producing these characters seem to segregate and recombine like any other factors. If the combination is such as will produce a pressure of
the head in a certain rate and intensity, various degrees of squareheadedness may result. If the head encounters little or no pressure the internodes may be more or less uniform, and if the growth of parts is unbalanced certain abnormalities of the spike may result.

Since a number of morphological factors are concerned in the production of equareheadedmes. logically it would be expected, and experimentally it would be found, that the segregates of a cross between a squarehead and a non-squarehead do not appear in a definite ratio but give a distribution approaching the normal curve of error.

\section*{EFFECTS OF NUTKITION}

In one of the preliminary experiments to determine the effect of nutrition under field conditions, seeds from a pure variety of a squarehead were grown at varying distances. In one case the seeds were drilled rather closely; in the second case they were planted 7.5 centimeters apart; in the third case they were planted is centimeters apart. The frequency distribution of squareheadedness of these three sets of plantings is given in table 1 . The set that was drilled in had a mean degree of squareheadedness of \(1.325 \pm 0.012\); the seeds planted 7.5 centimeters

TABLE 1. Variations of SQuareheadedness in Plants Grown at Varying Distances
(Variety; Giant Squarehead)
\begin{tabular}{|c|c|c|c|c|c|c|c|c|c|c|c|c|c|c|c|c|}
\hline & \multicolumn{13}{|c|}{Squareheadedness} & \multirow[t]{2}{*}{\[
\begin{aligned}
& \text { Number। } \\
& \text { of } \\
& \text { plants }
\end{aligned}
\]} & \multicolumn{2}{|r|}{\multirow[t]{2}{*}{Mean}} \\
\hline & 1.00 & 1.10 & 1.20 & 1.30 & 1.40 & 1.50 & 160 & 1.701 & 1.801 & 1.90 & 2.00 & 2.10 & 2.20 & & & \\
\hline Seeds drilled & 1 & 14 & 18 & 25 & 19 & 3 & . 10 & 2 & & 2 & & & & 94 & & \(325 \pm .012\) \\
\hline Seeds planted 7.5 cm a apart & .. & & 1 & & 8 & 3 & 10 & 7 & 3 & 3 & 2 & 4 & 1 & 42 & & \(1.678 \pm .025\) \\
\hline Seeds planted 15 cm apart & & 2 & 6 & & 2 & & 1 & & & & & & & 11 & & \(254 \pm 027\) \\
\hline
\end{tabular}
apart gave a higher mean, \(1.678 \pm 0.025\); and those planted i5 centimeters apart gave a mean of but \(1.254 \pm 0.027\).

The plants from the drilled rows made a fair growth but were inferior to those of the second set. The plants of the third set were mostly winterkilled, and such as survived had a poor stand with heads of varying length and degrees of development. The poor condition of the lastnamed was due to the wide distances between the plants, which made them unable to protect themselves from winter conditions. In the case of the other sets there was enough foliage developed during the fall for winter protection.

Dinregarding the thind set. : in afparent that the high degree of equareheadedness of the second set was due to the greater feeding allotted in these plants: for there wa- aho a corre-pmonding general developnent.

Another experiment was'made with potted plants growing under greenhouse conditions. It consisted of four sets of nine pots. One set was grown in a cool house, the second under moderate greenhouse conditions, the third in a damp chamber, and the fourth in a rather warm place. Each set consisted of triplicate pots containing, respectively, soils of a very poor sandy mixture, of fair fertility, and of higher fertility.

In the sets grown in cool and moderate temperature conditions, the pots containing poor soil produced heads of a low degree of squareheadedness, while the heads of plants grown in moderately fertile soils showed a higher degree of squarcheadedness. \({ }^{4}\)

There is no cuestion that in thene cane the hish degree wi - puareheadedness was produced by the fertility of the soil. That fertility in-
 Ohlmer (Igo8), and Meyer (Igog). Meyer found in addition that nitrogen was the causative factor, as neither calcium, potassium, nor phosphorus, alone or in combination, had any noticeable effect on the production of this character.

It is not so difficult to explain how high fertility increases squareheadedness, in the light of the causes of squareheadedness given in the preceding discussions. If rapid growth of the culm subjects the head to higher pressure, the spike takes the squarehead form. By increasing the nitrogen content of the soil, the rate of growth of the culm is accelerated and the tissues of the parts of the head are softened. The first of these conditions increases the pressure to which the head is subjected, and the second renders the head more sensitive to the effect of pressure. In the absence of sufficient nitrogenous food, the rate of growth is retarded and the parts of the head become fibrous. Due to the first condition sufficient pressure is not developed, and with the hardening of the tissues the spike offers a greater resistance to whatever pressure may the developed.

\footnotetext{
4The sets grown in the damp chamber and in the warm place did not do well.
}

The effect of the rate of growth on squareheadedness may lie determined also by ascertaining the degree of squareheadedness of the leading culm and of those that develop later. Practically in every case the leader, which is by far the most vigorously growing culm, has a higher desree of syuareheadednes than the others. Giten the smaller culms of squarehead plants will have vulgare-like heads.

In cases in which squareheading is inçreased by the rapid growth of the culm, the plants having longer spikes are more squareheaded than those with shorter spikes. The relative degree of squareheadedness of short and of long spikes of the same plant is shown in table 2. From fonn to six well-developed heads were measured from each plant in connection with another experment. Here the shortest and the longest of these, respectively, are shown. Ont of twelve cases taken at random there was but one case in which the short head had a higher derree of squareheadedness. The average of the summation of the differences was \(0.17 \pm 0.021\) in favor of the long heads.

TABLE 2. Differences in Degree of SQuareheadedness of Long and of Short Heads of the Same Plant
\begin{tabular}{|c|c|c|c|c|}
\hline \multicolumn{2}{|c|}{Short heads} & \multicolumn{3}{|c|}{Long heads} \\
\hline Length (centimeters) & Squareheadedness & Length (centimeters) & Squareheadedness & Difference in squareheadedness \\
\hline 100 & 1.23 & 13.5 & 1.47 & \(-0.24\) \\
\hline 11.6 & 1.41 & 13.2 & 1.51 & +0.10 \\
\hline 104 & 1.50 & 13.8 & 1.50 & 0.00 \\
\hline 103 & 1.24 & 12.0 & 1.51 & \(+0.27\) \\
\hline 10.7 & 121 & 140 & 1.37 & \(+0.16\) \\
\hline 105 & 124 & 13.6 & 1.48 & \(+0.24\) \\
\hline 8.0 & 1.39 & 10.5 & 1. 60 & +021 \\
\hline 7.6 & 1.31 & 10.0 & 1.65 & -0.34 \\
\hline 87 & 1.51 & 101 & 1.65 & +0.14 \\
\hline 8.4 & 1.33 & 11.2 & 1.57 & \(\div 0.24\) \\
\hline 93 & 139 & 12.5 & 1.58 & +0.19 \\
\hline 7.9 & 129 & 103 & 1.17 & -0.12 \\
\hline \multicolumn{5}{|c|}{Mean and average error} \\
\hline \(44 \pm 85\) & 124土 112 & \(\ldots\) &  & \(-117 \pm 021\) \\
\hline
\end{tabular}

When vigor is induced by soil fertility, the plants with longer heads will be more squareheaded than the others; and when vigor is induced Ly crossing certain autgare forms, the long heads of each plant will be fround to be more of a squareheaded type. The increase of squareheadedness of the \(F_{1}\) plants, as shown later, will serve as examples.

If, on the other hand, -rpuareheaderlness in caused, not su much liy the
vigorous development of the culm, but by the failure of the sheath to unfold at the proper time, then the plant, that are more spuarelieaded will be found to have shorter heads that the non-squarehead forms. Figures illustrating this type of -ytareheadedness are given in connection with the discussion of that subject.

Before concluding the discussion of the effects of nutrition, it may be well tomake a tew remathe recarime it effect on the density of the compactum form. As already mentioned, the density of this form is not the result of pressure. But by increasing the fertility of the soil it is possible to change markedly the degree of density. Four heads from each of two compactum plants are shown in Plate LXVIII (lower). These two plants were grown in 4 -inch pots in a greenhouse. The soil was highly fertilized. The heads first developed (2, 4, and 5) were almost like rulgare, but as the season advanced, and more spikes began 10 head out, the heads became more and more dense. There was an interval of about four weeks between the time of heading-out of the first and of the last head. When the last spike headed out, the first one was almost ripe-that is to say, the nutrients in the plant or those in the soil were already used up; hence the spikes heading out later obtained very little food. From the difference in thickness of the culms of dense and lax heads an idea may be formed of the relative amount of nourishment obtained by the different heads.

This experiment was conducted under abnormal envirommental conditions, and it is not likely that variations as great as these will be found on plants growing under field conditions. But it points out the fact that increa-ed fertility in the ent temds the increace the leneth of the rachis internodes.

\section*{SUMIMARY}

The wheat plant during its development undergoes two more or less distinct periods of growth. In the first period the sheaths and the blades develop. In the second period the rate of growth of the sheaths diminishes and the culms begin to develop, and during this period the spike carried at the end of the terminal culm internode is pushed up through the enveloping sheath.

Squateheadednes is the combination of a mumber of character which
are produced by the shortening of the terminal rachis internodes. It is expressed by the cocfficient found by dividing the number of internodes in the terminal third of the rachis by the number of internodes in the central third.

Density is the shortening of all the rachis internodes. It is determined by dividing the length of the rachis in millimeters by the number of internodes.

There are numerous gradations of squareheadedness and of density.
Squareheadedness is the result of pressure developed by differential growth of culm and sheath.

Rapid growth of the culm, failure of the sheath to unfold, and ears with soft-spreading glumes, tend to increase squareheadedness.

Any factor, genetic or environmental, which affects principally the development of the above-named characters, will affect the degree of squareheadedness.

Fertility of the soil or access of the root system to sufficient available nitrogenous matter increases squareheadedness.

Density is purely an inherited character, but favorable growth conditions may somewhat increase average internode length. Under abnormally farorable or unfavorable conditions, the increase or the decrease of density even on the same plant may be considerable.

\section*{THE GENETICS OF SQUAREHEADEDNESS}

Investigations on the genetics of squareheadedness have given many confu-ing revits, chicfly becaus no definite standard have been used for measuring this character. There are many instances in which this word has been used for designating a moderate degree of density.

One of the earliest studies of the subject was made by Rimpau (IS9I), who crossed lax zulgare types with lax and moderately dense squareheads. The \(\mathrm{F}_{1}\) hybrids were intermediate and the types of the \(\mathrm{F}_{2}\) populations varied within respective parental ranges.

Von Rümker (1909) obtained from squarehead \(x\) zulgare crosses, \(\mathrm{F}_{2}\) populations, some of which yielded more and others fewer squareheads. The squareheads varied also in degree.

Nilsson-Ehle (19II) found the aulgare type to be dominant over the
squarehead form, The ratio of zulgare to squarehead was between \(3: 1\) and \(15: 1\).

Further work has been done, fint becanse of the different meanings given to the word squarehead it is not posible to compare the zesults with those that are here presented.

\section*{INHERITANCE OF SQUAREHEADEDNESS IN CROSSES WITHIN THE SATIVUM GROUP}

Squareheadedness is a quantitative character. Crosses in which it is insolved do not show a clear-cut segregation into mendelian chases and ratios. Phesological experiments have shown that the character is the result of the interaction of a number of growth factors, certain combinations of which canse the compacting of the teminal part of the spike. The character is very variable, for any environmental condition that affects these growth factors in one way or the other indirectly increase or decreases the degree of compactness of the terminal fart of the rachis.
Before considering the inheritance of the character in spuarehead \(x\) non-oquarehead crosses, it may the well willustrate the mode of inheri-

Table 3. Degree of Squareheadedness in Vut.gare x Vulgare Crosses

tance of spuareheadednes when ether the spuarehedds or the erne ete type（non－squareheads）are crossed among themselves．

The degree of squareheadednes in \(F_{1}\) and \(F_{z}\) gencrations of onf w wo \(^{\circ}\) A eutyare crosses is shown in table 3．The first three croses here eerets I3I5 a，I3I－8a，and I3I59a are between Mealy and lones Lonstiers： both of which have practically the same degree of spuarelneadednes．I It and I．12，respectively．The mean degree of squareheadedness of the \(F_{\text {a }}\) generation fluciuated around the mean－wi their［arental forms，fome， respectively，I．I2，I．IT，and I．II．

The fourth cross，I3Iララa，was made between plants of lower coeffi－ cient ；that of Pride of Genesee was I．OS and that of Dawson Golden Chaff was 0.98 ．The \(F_{2}\) generation from this cross were all non－square－ head－and had a mean squareheadedness of i（ow ，these al－u＝omewhat ati－ proaching the average of their parents．

The degree of squareheadedness in crosses between squareheads is shown in table 4．The \(\mathrm{F}_{2}\) of the first cross，I 3201 a ，has a range with－

TAELE 4．Deqree of Sguarehlatedxes in Squarehead x Squarehead Comese

in the squarehead classes with a mean of I．46．Compared with the averages of the parental forms－New Sonles，I．万I，and Giant Square－ head， 1.77 －the mean of the \(F_{2}\) is lower．The second cross，I3203a， has an \(F_{2}\) mean squareheadedness within the means of the parents．

There are two points of interest in connection with these two sets of crosses：first，as a rule，when iulgare forms are crossed among them－ selves or squareheads are crossed among themselves，the \(F_{2}\) generation

TABLE 5. Degree of Squareheadedness in Squarehead x Vulgare Crosses

consists of practically only oulgare or only squareheads, respectively; secondly, the mean of the \(\mathrm{F}_{2}\) generation approaches the average of the parental forms.

Kegarding the inheritance of symareheadednes amons squarehead \(x\) vulgare crosses, an idea can be obtained by comparing the \(\mathrm{F}_{2}\)-generation distributions with the parental distribution (table 5). Special attention might be called to the comparatively high degree of squareheadedness of the \(F_{1}\) generations; in most of the cases the \(F_{1}\) plants are almost as squareheaded as the squarehead parents. On a theoretical basis the means of the \(F_{1}\) would be expected to coincide with those of the \(F_{2}\). The departure here is too wide. This increase in squareheadedness of the \(\mathrm{F}_{2}\) is attributed both to heterosis and to greater care taken in growing and spacing the plants of this generation.

The \(\mathrm{F}_{2}\)-generation distribution, even when the number of individuals of which they are composed is considered, shows certain characteristics with respect to range of distribution and mean. If the \(\mathrm{F}_{2}\) distributions of squarehead and non-squarehead.crosses are compared with those of the crosses in which the parents were either both squareheads or both vulgare, it is seen that so far as the mode of inheritance is concerned there is no essential difference between them. The means of the \(\mathrm{F}_{2}\) generation plants shift toward on allaty from the mome squarebead clanes. but show a constant tendency to regress toward the means of the parental forms; and the range of the \(\mathrm{F}_{2}\) also spreads or contracts, depending on the extent to which the parents vary in degree of squareheadedness.

Although several of these crosses were carried through the \(F_{3}\) generation because of the similarities of the results, only two series are considered here to illustrate the behavior of the \(F_{2}\) plants in \(F_{3}\). The results of Giant Squarehead x Fultzo-Mediterranean (series i3I.40a) are shown in table 6, and those of Giant Squarehead \(x\) Mealy (series I3I35a) are shown in table 7.


SQU'areheadedness and Density in Wheat


That a segregation of forms is taking place is (fuite evident. The \(\mathrm{F}_{3}\) distributions shown in table 6 are arranged in the order in which they were planted, so that the differences may not be ascribed to envirommental conditions. Cultures 2.-4, \(25-3\), and \(25-4\) show notable differences in distribution and mean from the cultures growing next to them. Similarly, in table 7 , where the \(\mathrm{F}_{3}\) distributions are arranged according to the degree of squareheadedness of the \(\mathrm{F}_{2}\) plants, the difference in squareheadedness of cultures 19-17, 19-18, and 19-19, and many others, is to be noted. The progeny of plant ig-I\%, representing a line in which none of the plants were squareheads, grew immediately next to a row, 19-18, which produced only squareheads, thirty in all.

In order to show that the variations noted in the \(\mathrm{F}_{3}\) are not chance variations but are hereditary, the curves shown in figure 78 were plotted,


Fig. 78. retation of degree of squtareheadedness between fo parent plants AND THEIR PKOCENY
based on the \(\mathrm{F}_{3}\) data of series i3I40a (table 6). Tiwo sets of curves are shown in this figure. The dotted curve represents the degree of squareheadedness of the \(\mathrm{F}_{2}\) plants, and the solid curve represents, the means of the \(\mathrm{F}_{3}\) in the order of squareheadedness of the \(\mathrm{F}_{2}\) plants.

The inclination of the straight line fitted to the curve of the \(F_{3}\) progeny shows that in a general way the degree of squareheadedness of the progeny is determined by the degree of squareheadedness of the \(\mathrm{F}_{2}\) parent plants. A few words of explanation may be given to account for the constant rise and fall of this curve. Since the \(F_{3}\) figures represent, respectively, the means of approximately thirty individuals, there is no question that the are dependable, and the differences between the means of the \(\mathrm{F}_{3}\) plants show their comparative genotypic differences. Since the mean of the \(\mathrm{F}_{3}\) in material having a mode of inheritance such as this character fluctuates more or less around the degree of squareheadedness of the \(F_{2}\) fom planted, it may be assumed that, had the \(F_{2}\) plants been grown in the same year as and under conditions similar to those under which the \(\mathrm{F}_{3}\) were grown, the average degree of squareheadedness of the developed heads of the \(\mathrm{F}_{2}\) planted would have approached more nearly the mean of their progeny. Because of the large number of plants that had to be measured, it was not found practicable to measure several heads from each \(\mathrm{F}_{2}\) plant. A developed head was taken at random from each of the envelopes containing the heads of each plant, and was measured. Since different heads in a single plant vary greatly in - puateheatednces, the developed head evected at random does not necessarily show the true phenotype. Therefore, the zigzag nature of the \(F_{: 3}\) curve should not be ascribed to inconsistent variations, but to inability to ascertain the true degree of squareheadedness of the \(\mathrm{F}_{2}\) by a single measurement, or even more, of the heads of a single plant.

The difference between the slope of the straight line of the \(F_{2}\) curve and that of the \(F_{3}\) is due to two conditions. In the first place, while the \(F_{2}\) curve.represents individual plants of a single frequency distribution, the \(F_{s}\) curve represents the means of such distributions. In the second place, as the mean represents the average of all plants exhibiting variations, slight or extreme in either direction, the means of the \(F_{3}\) cannot vary as much as the individuals of the \(\mathrm{F}_{2}\) in which the extremes
 straight line of the \(F_{3}\) means could therefore not be expected to be as steep as that of the \(\mathrm{F}_{2}\)-generation plants.
squareheadednes is not pectitar the tax form commonly known as squarehead. It may be present or absent in other forms also. The comfothm forms wowally are squareheaded but there are alow many which are not. In the examples that follow it is shown that squareheadedues introduced by a club has the same effect on the forms of the succeding generations as does spuareheadedness introdtced by the ordinary lax squarehead.

The dense spuarchead of the compacto-capitatum form used in this study was that known as Dale Gioria (Plate LXVII, upper, 5). This form is very dense and has an aterame internode leneth varyine usually.
 low to high, as shown in table 8 . With very dense wheats such as Dale
 Cross
(Series 13173a, Dale Gloria x New Soules)


Gloria, there is a certain limit beyond which the internodes do not shorten further. This is particularly true when the terminal spikelets are fertile, for development of a spikelet and development of adjacent partsglumes, nodes, internodes--seem to go hand in hand. A slight increase in the length of the terminal internodes of heads having an internode length in this region as short as 0.8 to 1.2 millimeters, greatly lowers the degree of squareheadedness. This accounts for the presence of forms of a low degree of squareheadedness among Dale Gloria plants.

In a cross between Dale Gloria and New Soules (series I3I;3a, table 8 ), in which both parents were squareheaded, the \(F_{2}\) plants were found to be practically all in the squarehead classes. There were a few in the non-squarehead classes but these were all dense forms like Dale Gloria.

In a cross such as this, there is alon an independent segregation of densits in the \(F_{2}\) generation. Half of these \(F_{2}\) plants were carried through \(F\) The F forms were practically all squareheaded. In some cased there w.re a few non-squarehead= but these were in a very low proportion and probably were genotypically sguareheads. Is far as squareheadednest is concerned, the Dale Gloria \(x\) New Soules cross lehaved like any squarehead \(x\) squarehead cross, such as those shown in table +1 pase 855).

When Dale Gloria is crosed with z'ulyare forms (table 9), the distriTABLE 9. Inheritance of Squareheadedness in Vulgare \(x\) Compactum Crosess
(1321ła, Turkey x Dale Gloria; 1337a, Turkish Amber x Dale Gloria; 13172a, Mealy x Dale Gloria)

butions of loth the \(F_{1}\) and \(F_{2}\) generations show a marked resemblance to those profuced by squarehead a aulyare crosses as shown in table 5 (page SIf). The curves of the \(\mathrm{F}_{\mathrm{a}}\) in hoth instances range from the awlgave to the squarehead classes. In this case also Dale (iloria behatred ai a squarehead; and it is in reality a spuarehead. but in addition it happens to carry a density factor.

Two of the crosses in table 9 were carried through \(\mathrm{F}_{3}\). The results were similar to those already observed in tahles on and 7 page= RIS and 819). Using class I . 30 as an arhitrary line doviding mon-squarehead= from: squareheads, different \(F_{z}\) progenies produced non-squarehead, ans squareheads in different proportions and deerees. There were progenies which consi-ted of mothing but - puatehead, and othere which con-inter only of non-squareheads.

The question of the inheritance of syuareheadednces, and e-pecially that of the relationship of spuarelieaduches to cortain other characters, is considered further in the discussion of inheritance of density. The seneral fact o werved thus far resarimg the inleritance of squareheadedness among forms of the sativum group may be summarized as follows:

Squareheadedness is not a unit character, but is a resultant of a comEination of srowth character which parduce shorening of the terminal internodes. The \(\mathrm{F}_{2}\) and subsequent generations show segregation of Flants or line of symareheadednes ditinctly arsing in degree, but no definte ration are ofmerved among thene. The aberase desree of squareheadednes of the \(\mathrm{F}_{1}\)-rencration fant is wally much higher than the mean of the \(\mathrm{F}_{\mathrm{z}}\); in sme instance: in thin study it approached that of the squarehead parent. Apparent vigor due to heterosis, and the greater care usually given to \(\mathrm{F}_{1}\) plants, are considered to account for their variation.

Squareheads crossed among themselves or sativums crossed among themetre produce, sencrally -peahing, mily sumatheads on sutioums, as the cate may lee, of ransen and meann apmoathing thene of the parental forms. The crosses between squareheads and satioums show a wide range of variation, but the range and the mean of the \(\mathrm{F}_{2}\) are still determined by those of the parental forms.

Some of the clubs are squareheaded, and these behave as squareheads when crossed with other forms.
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RELATION OF THE DEGREE OF SQUARFHEADEDNESS OF THE, VULGARE
PARENT IN VULGARE X SQUARFHEAD CROSSES, TO TIIE SQUARE-
HEADEDNESS OF THEIK PROGENY

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The question as to the extent to which the parent plants influence the squareheadedness of their offspring may be determined by examining crosses in which one of the parents is the same in all crosses
 a number of series are shown in which different varieties of zulgare are crossed with Giant Squarehead. The \(F_{1}\) and \(F_{2}\) generations of these crosses are shown graphically in figure 79, in which the crosses are arranged according to the degree of squareheadedness of the vulgare parent.

Giant Squarenead Squarehead parent


Fig. 79. dfgref of sQuareheatienness of paresta and progeny when different vulcarf. forms were crossen with giant squarehean

The curves representing the means of both the \(\mathrm{F}_{1}\) and \(\mathrm{F}_{2}\) generations show intermediacy between the parental curves. The \(\mathrm{F}_{1}\) curve, as has already been mentioned, shows, as a rule, a higher degree of squareheadedness than the \(\mathrm{F}_{2}\). The \(\mathrm{F}_{2}\) curve shows a general rise of the mean more or less in proportion to that of the anlyare parental curve. The \(F_{1}\) curve, being based on measurements of very few plants, shows great irregularities although it follows the general lines of the \(F_{2}\) curve.

The curve to be considered is that formed by the \(\mathrm{F}_{2}\) means. Although this curve shows a general rise, it cannot be said that the means of the Fa increase or decrease directly in proportion with the means of the vulgare parents. There are other factors introduced by the zulgare parents which, combined with those contributed by the squarehead parent, tend to neutralize or accelerate the production of squareheadedness. For example, Fultzo-Mediterranean producel an \(F_{2}\) progeny of a very high degree of squareheadedness, and the corresponding rise of the. \(\mathrm{F}_{1}\) curve shows that this rise is significant. Nealy and Ninnesota 169 , although of a higher degree of squareheadedness, produced individuals of lower mean squareheadedness.

The influence of the zulgare parent on the squareheadedness of vulgore \(x\) squarehead-compact crosses is seen in table 9. In the crosses in this table the squarehead-compact parent is Dale Gloria. The vulgare parents, Turkey, Turkish Amber, and Mealy, have coefficients of 0.92, I.04, and I.14, respectively. The ranking of the \(\mathrm{F}_{2}\) means is in the same order, and coincidently the differences between them are in proportion with the differences existing between the zulyare parents.

Because crosses between squareheads produced only squareheads, and those between zulgare forms only zulgare forms, it cannot be concluded that the degree of squareheadedness of the offspring is entirely dependent on that of the parents, for the determining factor is not the degree of squareheadedness of the parental forms but the peculiar combination of factors introduced by each variety. The mean squareheadedness expressed by a variety is but the resultant of the effects of these factors.

\section*{I'ALSE DOMLINANCE OF SOLTAREHEADEDNESS}

Since the \(\mathrm{F}_{2}\) progeny of a zuldare \(x\) zulyare cross consists only of zulgare forms, theoretically one would expect such a cross to produce
only non－squareheads in \(F_{1}\) ．This has not always been the case．In iahle 3 ＇page 太la＇，giving the results of three Mealy \(x\) Iones Longherry crosses，the parental means are I．I4 and I．I2，averaging I．I3，and the means of the \(F_{2}\)－generation plants are I．I2，I．I7，and I．II，also averag－ ing I．I3．But the \(F_{1}\)－gencration plants in all three cases were di－tinctly squareheads，with means of I．35，I．50，and I．42，respectively．In this paper the figures for squareheadedness or for density of \(F_{1}\) plants rep－ resent the arerages of usually from five to ten heads of each plant．If the apparent degree of syuareheadedness of the \(F_{1}\) plants indicated their approximate genotypic make－up，it would be expected that the \(F_{2}\) dis－ tribution would range somewhat near the mean of the \(F_{1}\) plants．The F．plants in these three cases were all ableare，showing that the appear－ ance of \(F_{1}\) plants of a high degrce of squareheadedness in these cases did not show any type of dominance of squareheadedness．

These crosses may be contrasted with the Dawson Golden Chaff \(x\) Pride of Genesee cross（13177a），in which the \(\mathrm{F}_{1}\) plants were all vulgare， their mean approaching that of the \(\mathrm{F}_{2}\) ．

The appearance of squarcheads in the \(F_{1}\) sencrations of－cries 13158 E ， 13I－8a，and 13I59a，and the absence of such forms in 13I7うa，may be accounted for by the supposition that in the first three crosses the com－ linations of growth factor－contributed ley the parente were favorable to the production of afuareheads，while thoee contributed by the fatemt in series I3I㘯a were not favorable．

The presence of squareheads in \(F_{1}\) and their total absence in \(F_{2}\) in series 13158a，13178a，and 13I59a，may be regarded as cases of false dominance．

RELATION OF WIDTII OF CULM TO SQUAREHEADEDNESS
The part of the culm below the base of the spike tends to be wider in spuareheads than in head of uniform internode length．The data here considered were taken on the progeny of a lax squarehead \(x\) vulgare cross，which was not as favorable a material for the study of this charac－ ter as would have heen some other crosses in which a more intense squarehead was used．The results，however，were satisfactory enough to illustrate the degree of correlation between squareheadedness and width of culm．

In taking measurements for squareheadedness of the \(\mathrm{F}_{3}\) generation of series 13135 a , the width of the culms of some 260 heads was measured also. The measurement was taken about 2 centimeters below the basal rachis internode. The resulting data are shown in correlation form in table for. The cosfficient of correlation in this distribution in \(0.4^{10} 5 \pm\) 0.033 , which is significant.
table 10. Corrilation between Diameter of Culm and Squareheadedness (Series 13135a, Giant Squarehead x Mealy)

Degree of squareheadedness


The writer considers this correlation as not due to any form of linkage but an a dircet result of somareheadedne... which is caned in turn loy the shortening, or rather hampering of the deselopment, of the terminal part of the apike due to prenure producel ber rapid development of the culm internole immediately at the hase of the head, and to the failure of the sheath enveloping the spike to grow at a corresponding rate. When the longitudinal growth of the culm internode is checked or retarded, growth naturally takes place in other directions, often causing a thickening of the culm wall. A similar thickening of the wall of the culm occurs also in non-squarehead dense plants and more pronouncedly in squarehead dense plants. In these cases, however, the thickening seems to be due to the presence of the density factor, which shortens the culm internode length as well as the length of the rachis internode. Due to dwarfing, the plant cannot grow in height. The food produced constantly by the plant is stored partly in the culm, thus increasing the thickness of its walls.

The club wheat owes its ability to stand erect in the presence of strong winds to the presence of the density factor, which, as mentioned above, produces thickening of the culm and incidentally of other parts. This factor, which is later considered at length, causes the shortening of the culm also, without affecting the number of nodes. The shortening of the culm internodes increases the number of nodes to each unit of length, thereby giving the culn added -trength; moreoner, the shortening of the culm lowers the leverage. These three conditions, direct or indirect results of the presence of the density factor, are the chief causes of the nou-lodging quality of dense wheat.

INHERITANCE OF SQUAREIIEADEIDEESS IN SPELT X SATIVUM CROSSES
In crosses wherein the spelt character has been introduced, the curve of the \(\mathrm{F}_{2}\) generation is very distinctly skewed near the extreme of the range on the side of the non-squarehead classes (table II). True spelts

Table 11. Squareheadedness in Spelt x Sativum Crosses

differ in their ability to carry the factors producing squareheadedness. It cannot lee determined from the appearance of the -pelt plants whether or not they carry the squareheading factors, as the spelt character acts as an inhibitor for squareheadedness. In fact, the presence of a large number of individuals in the non-squarehead classes of spelt \(x\) satioum crosses is the result of the presence of a large number of spelts, which, although carting the -quareheadednes factors were themelves nom-
unareheads hecanse they carried aho the -pelt factur, which, as stated above, acts as an inhititor for the squarehead character.

Five crosses between spelts and satizums are shown in table 11. \({ }^{5}\)
 Spelt, produced \(F_{2}\) generations composed of non-squareheads only. When this same spelt furm wat crosed with biant Squarehead isernes 13216a), a number of squareheads were produced in \(\mathrm{F}_{2}\). The White Socha Dale ciloria crom 1132550 alow howed a fair mumber uf sumareheads (Plate LXVII, upper, 6 to I2).

From these results it is seen that the White Spelt does not carry the necessary factors for squareheadedness, since when it was crossed with : mhtare it producel no squarehead. Siluatheads appeared umly when a squarehead form was used as the sativum parent.

These four crosses were carried through \(\mathrm{F}_{3}\). The first two crosses produced practically nu -ymarehead-; a few were obtained, but the were not tested to ascertain their stability. The remaining crosses produced \(F_{3}\) progeny which were composed of forms of various degrees of squareheadedness. Since the spelt factor acted as an inhibitor, the spelts of the \(\mathrm{F}_{3}\) showed no squareheadedness. The non-spelts produced curves similar to those shown in tables 6 and 7 .

That there was no so-called repulsion between the spelt and squareheadedness factors was evident from the reappearance of squareheads among the progeny of some \(\mathrm{F}_{2}\) spelts, and from the absence of squareheads among the offspring of other \(\mathrm{F}_{2}\) spelts.

Another spelt form, known as Black Bearded Spelt, when crossed with a zulgare, Jones Longbery (series 3085a), produced a large number of oquathead. Thme form- were mere intensely - guate heated than those produced by the White Spelt x Giant Squarehead cross (I3216a).

Alout ten spelt x vulyare crosses, with Black Bearded Spelt as one of the parents, were examined by the writer, and in every case there were a large number of squareheads in the progeny, most of which were semi-dense.

The progeny of cross 3085 a were not carried through \(F_{3}\), but another

\footnotetext{
\({ }^{5}\) The F 2 segregations, of these crosses were in the proportion of 3 spelts or speltlike forms to 1 sanvum.
}

Crow betheen this same Black Bearied sifelt and at athore showed that most of the squareheads bred true.

If it is recalled that some of the \(F_{2}\) spelts of the squarehead \(x\) White Spelt cross produced squareheads in F while others that were thenotypically like the former did not, it will not be difficult to understand how the Black Bearded spelt conld have producel different rewhe from there of the White Spelt. It appears. from the ee examples, that spelts may carry the squareheadedness factors the same as do squarehead sativums themselves, but due to the presence of the spelt factor, which acts as an inhibitor, such spelts do not appear squareheaded.

This leads to the consideration of another condition. Since the presence of squareheadedness cannot be detected without a genetic analysis, one may come across a spelt form which, crossed with infgarc, may sometimes yield squareheads and sometimes not. Either such a spelt is heterozygous with respect to squareheadedness, or the variety to which it belongs has not been stabilized with respect to this character. As the investigator is gutided by apparent characters in purifying a line or in calling it a pure line, he cannot detect the segregation of non-detectable factors which is going on within his selected line.

EFFECT OF THE SPELT FACTOR ON SQUAREIIEADEDNESS
In the discussion of the inheritance of squareheadedness in spelt \(x\) saituru crosses, it was mentioned that the spelt factor inhibited square-

TabLE 12. Relation of SQuareheadedness to Spelting (Series 13216a, Giant Squarchead x White Spelt)

Degree of squareheadedness
\begin{tabular}{|c|c|c|c|c|c|c|c|c|c|c|c|c|c|c|c|c|}
\hline & \[
\begin{aligned}
& \bar{\infty} \\
& =
\end{aligned}
\] & \[
\begin{aligned}
& \hat{0} \\
& =
\end{aligned}
\] & \[
\begin{aligned}
& \bar{\delta} \\
& =
\end{aligned}
\] & \[
\begin{aligned}
& \hat{S} \\
& =
\end{aligned}
\] & \[
\bar{\xi}
\] & e & \[
\Xi
\] & "- & \[
\overline{9}
\] & \[
\bar{i}
\] & \[
\overline{\bar{x}}
\] & 约 & 三 & \[
\begin{gathered}
\text { K } \\
-1
\end{gathered}
\] & \[
\overline{20}
\] & : \\
\hline 1 & 1 & 3 & 3 & 3 & 2 & & & & & & & & & & & \\
\hline er 2 & 1 & 2 & 1 & 1 & & 2 & & & & & & & & & & \\
\hline . \({ }^{\text {E }}\) & 1 & & 1 & 1 & 2 & 2 & & & & & & & & & & \\
\hline \(\stackrel{1}{c}\) & & 1 & 3 & 1 & 1 & & & & & & & & & & & \\
\hline c, 5 & & 2 & 2 & 1 & 3 & & 1 & & & & & & & & & \\
\hline \(\bigcirc 6\) & & 1 & 1 & : & \(\cdots\) & & & & & & & & & & & \\
\hline \(\stackrel{0}{0} 7\) & & 1 & & & 1 & & & 1 & & & & & & & & \\
\hline 80 & & 2 & & 1 & 2 & & & & & 1 & & & & & & \\
\hline ๑ 9 & & & 1 & & 1 & 1 & 2 & 1 & 1 & & 1 & & 2 & 1 & & 1 \\
\hline 10 & & & & & & & & & & & 2 & & & 2 & & \\
\hline
\end{tabular}

headedness. The object here is to show to what extent this inhibition takes place. To illustrate this, three types of crosses are used. The first is the Giant Squarehead \(x\) White Spelt cross (table 12) ; the second in Whate spelt a Dale Ciloria (table 1 : 1 : and the thed in Iblack Bearded Spelt x Jones Longberry (table I4). In the first two crosses, squareheadedtexs in intrombed by a lax -quarchead and a denee squarehead, respectively, and in the last cross it is introduced by the spelt.

The \(\mathrm{F}_{2}\) plants in these tables are classified into ten arbitrary classes of spelting. The spelts in such crosses exhibit the spelt character in various degrees. Those showing it in an intense form are placed in class 1 ; classes \(2,3,4\), and so on up to 9 , show various degrees of dilution of the character; the plants in class 10 are all sativum forms, and lack the spelt altogether. \({ }^{6}\)

The distribution of the plants with re-pect to -pelting and suruareheadedness, as shown in tables 12 to 14 , seems to depend to a great extent on the types used as parents. Different spelts crossed with different sationms show different modifications. All crosses, however, show the following semeral di-tribution: Sicelt- of clase- I to 5 inclu-ise arrans themselves more or less within the inn-apuarehead clanses o. © on to I . I 5. There is a slight temdency in spelt clases 3 to 5 to mowe the mean toward 1.15. As the classes approach the sationm type, the shifting of the mean toward the more equareheaded clasen is accelerated in sermetrical proportion.

The spelt factor, then, does not prevent squareheadedness altogether, but plants that are heterozygous spelts or those that carry modifiers tending to dilute this spelt character are very much more likely to be qquartheated than plants that are intensely spelted. squareheadedness in spelts, however, is always of a low degree.

\section*{INHERITANCE OF SQUAREHEADEDNESS IN SPECIFIC CROSSES}

It ha- already heen shown (fage is crossed with a sativum of a uniform rachis internode length, an appreciable number of squareheaded forms appear in the \(F_{2}\). Squareheads may be ubtamed also when a culyure is crosed with other species. In table 15 the progeny of a number of interspecific crosses are classified into two gronps, the first containing forms reombling the sutioum type and the second including all the other forms. Their degree of squareheadedness is represented in the usual manner.

In all series a large proportion of the sationm or sativum-like forms were squareheads. The segregate lefonging to other species were practically all non-squareheads.

\footnotetext{
\({ }^{6}\) The genetics of the spelt character with reference to the crosses under consideration has been fully discussed in a recent paper (Leighty and Boshnakian, 1921).
}

TABLE 15. Degree of Squareheadedness in Species Crosses. Fz Data
```

3032 Salt Lake Club x Kahle (Trilicum durum)
3034 Gharnovka (T. durum) x Black Bearded Spelt (T. spelta)
1312a Poole (T. vulgare) x Alaska (T, turgidum)
1328a Satisfaction (T. vulgare) x Alaska (T. turgidum)
1360a Jones Paris Prize (T. vulgare) x Kubanka (T. durum)

```

*Series 3034 did not produce true sativums in the \(\mathrm{F}_{2}\).
t 2.00 or over.
Squareheadedness is not confined, however, to the satioum form. Other species, with the exception of the wild wheat, may show this character, particularly the eceresate- uf the dumum, the furndum, and thr \(f\) hmionn tyes and eren the diciovme wpe. But spluareheadednew in thene forms in wif low dese and is comparatively rate (Plate LAVII, lower).

The analysis of the \(F_{2}\) forms in specific crosses presents a difficult task because a large number of specific forms appear, of which some are develoned and others are bery medioce or sterile with different tendencies toward squareheadedness.

\section*{SUMMARY}

Syuarchedednes, heme the re-ult wit ambination of growth characters, shows a complex mode of inheritance. Simple mendelian segregations were not obtained in these experiments.

In the \(\mathrm{F}_{3}\) generation there were obtained plants of different degrees of syuareheadednes, ranging from foms which were distinctly enmpacted at the tip to forms which were denser near the base of the ear.

As a rule the range wi variation in \(F\) : depended on the differences hetween the extremes of the parental ranges. The means of the \(F_{2}\) approached the parental means.

The \(F_{1}\) generation unually had a higher coefficient than the praventai mean. Some \(F_{1}\) progeny of two non-squarehead parents were eren distinctly squareheads, but in \(F_{\text {a }}\) none of these forms were obtamed. These variations from normal expectations are ascribed to heterosis and th greater feeding area.

A coefficient of correlation of \(0.465 \pm 0.033\) was found between width of culms and squareheadedness.

The purely spelt forms were found not to be affected by the factors producing squareleadednes. The more the spelts appoached the sativum type, the more they were found to be affected. Speltoid forms did carry these factors, as is shown by the fact that among their satioum progeny a large number of squareheads of varying intensities were found.

Certain spelts when crossed with a vulgare form will produce a large number of squareheads in \(\mathrm{F}_{2}\). Others will produce only non-squareheaded sativum forms.
 other wheat species.

\section*{THE GENETICS OF DENSITY}

The discussion in the following pages deals with the genetics of compactness of the dense forms of wheat, and especially of Triticum compactum, the club wheat. The name club refleat seems to have been originally given to the squareheads, but at present it is applied almost exclusively to the compactum form.

Tine index of compactness used in this paper is the average rachis internode length. The index is found by dividing the length of the rachis in millimeters by the number of rachis internodes. The denser or more compact the head, the shorter is the length of the rachis internode.

Although there are numerous grades of compactness, the rachis internode length of what is usually called a club wheat does not exceed 2.25 millimeters. The mean density of the club used in the studies presented in this paper was about 1.4 millimeters.

INHERITANCE OF DENSITY IN CROSEES BETWEEN TRITICUM COMPACTUM AND OTHER FORAS OF TIIE SATIVUAI GROUP
The studies of density herein discussed were made on crosses be:ween Dale (Bloria (Plate L.NTII, upher, 5 ) and a number of lax forms consisting of both squarehead and vulgare types. The mean density of ihe Dale Ciloria parent wa- 1.41 millimeter-; the mean- of the las parents were in the neighborhood of 4.50 millimeters.

The \(F_{1}\) hybrids were all dense, but not quite as dense as Dale Gloria (table 16). They varied from I 80 to 2.40 , depending on the cross.

TAble 16. Secregation of Density in Crosses between Dale Gloria (Compactum) and Lax Forms
13174a Extra Early Windsor x Dale Gloria
13173a New Soules : Dale Gloria
13172a Mealy x Dale Gloria
13214a Turkey x Dale Gloria
13215a Seneca Chief x Dale Gloria
1337a Turkish Amber x Dale Gloria
13213a Red Wave x Dale Gloria


\footnotetext{
*Tbis figure represents the arithmetical average, not the mean of the frequency distribution.
}

The density curves of the \(\mathrm{F}_{2}\)-generation plants, unlike those of the aguarehoad \(x\) non-spuarehead crosses, were all discontinuous. consisting of two well-defined curves. In each case the curve of the dense classes contained about three times as many individuals as that of the lax classes. The proportions of these forms are given in table 17. In five cases the

TABLE 17. Proportions of Dense and Lax Segregates of the Fr Generations of Crosses between Dale Gloria (Compactum) and other Lax Forms
\begin{tabular}{|c|c|c|c|c|c|c|c|c|c|c|c|}
\hline \multirow{2}{*}{Series} & \multicolumn{2}{|l|}{Dense forms} & \multicolumn{2}{|l|}{Lax forms} & \multirow{2}{*}{Deviation} & \multirow{2}{*}{Probable ertor} & \multirow{2}{*}{\begin{tabular}{l}
Dev. \\
P. E.
\end{tabular}} & \multirow[b]{2}{*}{Mean of dense plants} & \multirow[b]{2}{*}{\[
\begin{aligned}
& \text { Mean of } \\
& \text { lax } \\
& \text { plants }
\end{aligned}
\]} & \multirow{2}{*}{Mean of all plants} & \multirow[t]{2}{*}{Total number of individuals} \\
\hline & Number obtained & Number expected & Number obtained & Number expected & & & & & & & \\
\hline 131743 & 40 & 45.7 & 21 & 15.2 & \(-+5.8\) & \(\pm 2.28\) & 2.54 & 1.88 & 3.95 & 2.59 & 61 \\
\hline 131733 & 39 & 36.0 & 9 & 12.0 & +-3.0 & \(\pm 2.02\) & 1.48 & 1.74 & 3.74 & 2.33 & 48 \\
\hline 13172a & 55 & 57.0 & 21 & 19.0 & \(\cdots+2.0\) & \(\pm 2.55\) & 0.78 & 2.11 & 4.04 & 2.64 & 76 \\
\hline 13214a & 54 & 50.2 & 13 & 16.7 & \(+-3.7\) & \(\pm 2.39\) & 1.55 & 2.30 & 4.51 & 2.74 & 67 \\
\hline 13215 a & 54 & 50.2 & 13 & 16.7 & +-3.7 & \(\pm 2.39\) & 1.55 & 1.98 & 4.12 & 2.40 & 67 \\
\hline 1337 a & 43 & 48.7 & 22 & 16.2 & \(-+5.8\) & \(\pm 2.36\) & 2.46 & 2.34 & 4.40 & 3.04 & 65 \\
\hline 13213a & 47 & 45.7 & 14 & 15.2 & +-1.2 & \(\pm 2.28\) & 0.53 & 2.33 & 4.67 & 2.87 & 61 \\
\hline Total & 832 & 333.7 & 113 & 111.2 & - +1. \(¢\) & \(\pm 6.16\) & 0.29 & 209 & 4.20 & 266 & 445 \\
\hline
\end{tabular}
deviations from the calculated ratios exceeded somewhat their protiable errors, and in two cases the deviations were well within the probable errors. Summing up the results of these seven crosses, of 445 plants obtained in the \(\mathrm{F}_{2}\) generation 332 were dense and II3 were lax; the deviation from the calculated proportions on the \(3: 1\) basis was -1.8 , which is about one-third of its probable error. These results show that so far as these crosses are concerned the \(F_{2}\) plants segregate into dense and lax forms in 3 :I ratio, the density being dominant.

Four of these crosses, of which two were with squareheads and the other two with vulgare, were carried through the \(\mathrm{F}_{3}\) generation in order to test whether the assumption of the presence of one factor was correct. The results obtained are condensed and given in table i8. Of the \(125 \mathrm{~F}_{2}\) plants tested, 30 were homozygous dense, 67 produced both dense and lax forms, and 28 were homozygous lax. These figures, compared with the calculated proportions-31.2 \(\pm 3.3,62.5 \pm 3.9\), and \(31.2 \pm 3 \cdot 3\), respectively-show a very close agreement with the \(1: 2: 1\) ratio. The propurtion of plant- whtaned in cach cros taken eparately agrees als, with the thereretical expectancy, the larse-t departure leing but 25 per cent more than its probable error.
T'ABLe 18. Proportions of Homozygous Dfn-1, Hfterozycits Denar., and Humozygous Iat Plants
Homozygous dense forms
\[
\|
\] Genfration
Homozygous la
\[
\left\lvert\, \begin{gathered}
\text { Total } \\
\text { numbler }
\end{gathered}\right.
\]



The behavior of each individual \(F_{2}\) plant of the four crosses under consideration is shown respectively in tables \(19,20,21\), and 22. Of the

TABLE 19. Behayror of \(\mathrm{F}_{2}\) Plants in \(\mathrm{F}_{3}\). Series 13172a, Mealy (Vulgare) x Dale Gloria
\begin{tabular}{|c|c|c|c|c|c|c|c|c|c|c|c|}
\hline & & tan & Mean & Dense & plants & Lax p & plant \(=\) & & & & \\
\hline pedistee & density & \(\left\lvert\, \begin{aligned} & \text { or dense } \\ & \text { Fs plants }\end{aligned}\right.\) & F3 plants & Number obtained & Number calculated & Number obtained & \[
\left\lvert\, \begin{gathered}
\text { Number } \\
\text { calcu- } \\
\text { lated }
\end{gathered}\right.
\] & F3 plan & & & \\
\hline 30-11 & 1.50 & 1.35 & \(\ldots\) & 29 & & \(\cdots\) & & 29 & & & \\
\hline \(30-21\) & 1.33 & 1.66 & ... & 22 & \(\cdots\) & . & \(\ldots\) & \({ }^{22}\) & & & \\
\hline 29-9 & 1.57 & 1.18 & \(\cdots\) & \({ }^{6}\) & . & & . & \({ }_{20}^{6}\) & & & \\
\hline 30.2 & 1.71 & 1.37 & . & 22 & & & \(\ldots\) & 22 & & & \\
\hline \(30-14\) & 1.73 & 1. 52 & \(\ldots\) & 24 & .... & & \(\ldots\) & \(2 \pm\) & & & \\
\hline \(30-5\) & 1.66 & 1.54 & .... & 24 & \(\ldots\) & & \(\ldots\) & 24 & \(\ldots\) & & \\
\hline 30-13 & 1.81 & 1.43 & & 28 & & & \(\ldots\) & 28 & & & \\
\hline 29-4 & 1.89 & 1.71 & .... & 18 & & & & 18 & & & \\
\hline 30-7 & \(1.8 \pm\) & 1.47 & 3.08 & 24 & 23.5 & - & 7.5 & 30 & +-1.5 & \(\pm 1.63\) & 0 92 \\
\hline 30-6 & 1.85 & 1. 72 & 3.41 & 33 & 32.2 & 10 & 10.7 & 43 & +-0.7 & \(\pm 1.92\) & 0.36 \\
\hline 30-9 & 1.90 & 1.52 & 3.01 & 34 & 33.7 & 11 & 11.2 & 43 & +-0.2 & \(\pm 1.96\) & 1.00 \\
\hline 30-3 & 1.95 & 1.69 & 3.56 & 31 & 30.7 & 10 & 10.2 & 41 & +-0.2 & \(\pm 1.87\) & 0.11 \\
\hline 29-6 & 2.00 & 1.44 & 3.06 & 18 & 18.7 & 7 & 6.2 & 25 & - +0.8 & \(\pm 1.46\) & 0.55 \\
\hline 30-19 & 2.00 & 1.65 & 3.00 & 23 & 21.0 & 5 & 7.0 & 28 & +-2.0 & \(\pm 1.55\) & 1. 29 \\
\hline 29-10 & 2.06 & 1.80 & 3.70 & 25 & 24.7 & 8 & 8.2 & 33 & +-0.2 & \(\pm 1.68\) & 0.12 \\
\hline 30-3 & 2.10 & 1.67 & 3.53 & 28 & 27.0 & 8 & 9.0 & 36 & +-1.0 & \(\pm 1.78\) & 0.56 \\
\hline 29-3 & 2.11 & 2.08 & 3.30 & 19 & 15.7 & 2 & 52 & 21 & +-3.2 & \(\pm 1.34\) & 2.39 \\
\hline 30-13 & 2.14 & 1.70 & 3.33 & 34 & 36.0 & 14 & 12.0 & 43 & -+2.0 & \(\pm 2.02\) & 099 \\
\hline 29.12 & 2.19 & 2.04 & 4.03 & 22 & 18.7 & 3 & 62 & 25 & +-3.2 & \(\pm 1.46\) & 2.19 \\
\hline 30-20 & 2.25 & 1.83 & 3.64 & 22 & 24.7 & 11 & 8.2 & 33 & - + 2.8 & \(\pm 1.68\) & 1.67 \\
\hline 29-1 & 2.39 & 1.90 & 3.15 & 18 & 16.5 & 4 & 55 & 22 & +-1.5 & \(\pm 1.37\) & 1.03 \\
\hline \(23-11\) & 2.42 & 1. i2 & 3.69 & 30 & 29.2 & 9 & 9.7 & 39 & +-0.7 & \(\pm 1.82\) & 0.3y \\
\hline 30-10 & 2.43 & 1.93 & 3.52 & 32 & 27.7 & & 9.2 & 37 & +-4.2 & \(\pm 1.78\) & 2.36 \\
\hline 30-1 & 2.50 & 2.22 & 4.23 & 37 & 35.2 & 10 & 11.7 & 47 & +-1.7 & \(\pm 2.00\) & 0.85 \\
\hline \(30-15\) & 2.52 & 1.85 & 3.47 & 28 & 25.5 & & 8.5 & 34 & +-2.5 & \(\pm 1.70\) & 1.47 \\
\hline 29-2 & 2. 53 & 2.00 & 3.64 & 19 & 19.5 & , & 6.5 & 26 & -+ 0.5 & \(\pm 1.49\) & 0.33 \\
\hline \(30-4\) & 258 & 1.81 & 3.56 & 27 & 23.5 & 11 & 9.5 & 38 & -+1.5 & \(\pm 1.80\) & 0.83 \\
\hline 29-14 & 2.84 & 1.78 & 426 & 13 & 13.5 & 5 & 4.5 & 18 & 1-+0.5 & \(\pm 1.24\) & 0.40 \\
\hline \multicolumn{3}{|l|}{Total heterozygous plants} & & 517 & 501.7 & 152 & 167.2 & 669 & |+-15.2| & \(\pm 7.551\) & 201 \\
\hline 30-16 & 3.28 & & 3.39 & & & 16 & .... & 16 & ... & \(\cdots\) & \\
\hline 30-12 & 3.55 & .... & 3.36 & \(\because\) & \(\ldots\) & 23 & \(\ldots\) & 23 & & .... & \\
\hline \(30-22\) & 3.75 & \(\ldots\). & 3.12 & \(\cdots\) & \(\cdots\) & 20 & \(\ldots\) & 20 & & & \\
\hline \({ }_{29-13}^{30-17}\) & 3.84
4.16 & \(\ldots\) & 3.43
3.88 & . & \(\ldots\) & 178 & .. & 18
20 & & & \\
\hline \({ }_{29-5}^{29-13}\) & 4.10
4.27 & & 3.88
8.65 & & & 17 & & 17 & & & \\
\hline 29-7 & 4.4 & & 4.08 & & & 5 & & 5 & & & \\
\hline \multicolumn{2}{|l|}{Total Fs plants.} & 1 & 1 & & & & & 980 & & & \\
\hline
\end{tabular}

609 heterozygous plants obtained in series 131 ²a (table 19). 517 were dense and 152 were lax, the deviation from the calculated ratio being about twice its probable error. In series i3I73a (table 20), the ratio between the probable error and the deviation from expectation is \(1: 5.22\), which is rather high. In series I3I万4a (table 2I) this ratio is I:5.02, and in series \(13214 a\) (table 22) it is only \(1: 0.46\). All these results, in spite of the difference hetween oborred an! calculated ration which

TABLE 20. Behavion of \(F_{2}\) Plants in \(F_{3}\). Series 13174a, New Soules (Capitatem) x Dale Gloria
\begin{tabular}{|c|c|c|c|c|c|c|c|c|c|}
\hline \multirow{2}{*}{\[
\stackrel{\mathrm{F}_{2}}{\text { pedigree }}
\]} & \multirow{2}{*}{\[
\underset{\text { density }}{\mathrm{F}_{2}}
\]} & \multicolumn{2}{|l|}{Dense plants} & \multicolumn{2}{|r|}{Lax plants} & \multirow{2}{*}{\begin{tabular}{l}
Total \\
F3 plants
\end{tabular}} & \multirow{2}{*}{Deviation} & \multirow{2}{*}{Probable error} & \multirow{2}{*}{\[
\frac{\text { Dev. }}{\text { P. E. }}
\]} \\
\hline & & Number obtained & Number calculated & Number obtained & Number calculated & & & & \\
\hline \[
\begin{aligned}
& 32-7 \\
& 31-11 \\
& 32-6 \\
& 31-3 \\
& 31-9 \\
& 328
\end{aligned}
\] & \[
\begin{aligned}
& 1.31 \\
& 1.52 \\
& 1.52 \\
& 1.55 \\
& 1.55 \\
& 1.62
\end{aligned}
\] & \[
\begin{array}{r}
37 \\
23 \\
-\quad 60 \\
40 \\
35 \\
41
\end{array}
\] & \(\cdots\) & \(\because\) & ....
\(\cdots\)
\(\cdots\) & \[
\begin{aligned}
& 37 \\
& 23 \\
& 60 \\
& 40 \\
& 35 \\
& 41
\end{aligned}
\] & \(\ldots\) & \(\ldots\)
\(\ldots\) & \\
\hline 31-8 & 164 & 24 & 278 & 13 & 9.2 & 37 & \(-+3.8\) & \(\pm 1.78\) & \({ }_{2} 13\) \\
\hline 32-13 & 165 & 52 & 58.5 & 26 & 195 & 78 & - 6.5 & \(\pm 2.58\) & 252 \\
\hline 32-14 & 168 & 51 & 34.0 & 21 & 180 & 72 & - 3.0 & \(\pm 2.48\) & 1.21 \\
\hline \(31-7\)
\(32-15\) & 1.71 & \({ }_{9}^{29}\) & 315
998 & 13
40 & 105
332 & 133 & -+2.5 & \(\pm 1.89\)
\(\pm .37\) & 1.32 \\
\hline \(32-9\) & 1.78 & 67 & 67.5 & 23 & 225 & \(90^{\circ}\) & -+ 05 & \(\pm 2.77\) & 0.18 \\
\hline 32-16 & 1.81 & 59 & 63.0 & 25 & 21.0 & 84 & -+4.0 & \(\pm 2.68\) & 149 \\
\hline 31-12 & 1.88 & 35 & 323 & 8 & 107 & 43 & +-2.7 & \(\pm 192\) & 141 \\
\hline 32-5 & 1.91 & 46 & 465 & & 155 & 62 & -+ 0.5 & \(\pm 2.30\) & 0.22 \\
\hline 32-10 & 1.95 & 62 & 63.0 & 22 & 21.0 & 84 & -+1.0 & \(\pm 268\) & 0.37 \\
\hline 31-1 & 1.95 & 30 & 43.5 & 28 & 145 & 58 & \(-+13.5\) & \(\pm 2.22\) & 608 \\
\hline \(32-2\) & 2.00 & 42 & 39.0 & 10 & 130 & 52 & +-3.0 & \(\pm 2.11\) & 1.42 \\
\hline 31-2 & 2.06 & 40 & 398 & 13 & 13.2 & 53 & +-02 & \(\pm 2.13\) & 0.09 \\
\hline \(31-4\)
\(32-12\) & 2.09
2.15 & 30
83 & 33.0
87 & 14 & 11.0 & 44 & -+3.0 & \(\pm 1.94\) & 1.55 \\
\hline 32-12 & 2.15 & 83
62 & 87.8
698 & 34
31
31 & 292
23
2 & 117
93 & + +4.8
-+7.8 & \(\pm 3.16\)
\(\pm 2.82\) & 1.52
2.76 \\
\hline \multicolumn{2}{|l|}{Total heterozy gous plants} & s 805 & 856.5 & \(\overline{3} 37\) & 2855 & 1.142 & \(-+51.5\) & \(\pm 987\) & 522 \\
\hline 31-10 & 3.40 & & & 37 & & 37 & & & \\
\hline \(31-6\) & 3.43 & \(\cdots\) & \(\ldots\) & 21 & \(\ldots\) & 21 & & & \\
\hline 32-4 & 3.45
3.80 & \(\because\) & \(\ldots\) & 56
69 & \(\ldots\) & 56
69 & & & \\
\hline 32-1 & 3.87 & \(\cdots\) & & 31 & & \({ }_{31}\) & & & \\
\hline \(31-5\) & 3.95 & & & 33 & & 33 & & & \\
\hline 31-13 & 4.19 & & & 24 & & 24 & & & \\
\hline 32-11 & 4.28 & & & 71 & & 71 & & & \\
\hline \multicolumn{2}{|l|}{Intal F3 plants} & & & & & 1.720 & & & \\
\hline
\end{tabular}
 only one factor is involved in the production of density.

Referring to tables 19, 20, 21, and 22, it will be noticed that while in
 rather closely, in series I3I73a and I3I74a an excess of lax forms was recorded in practically every case. In series 13I73a, out of 16 heter"rgons plants tered all hat three yielded an excess of lax forms, while in series I3Iク4a all but one plant yielded an excess of lax forms.

When wheat plant are grown clo-ely together, the routs weasionally tond to intertwine, and males the head- how variations of sesregating srose characters it in wot unally posible to determine whether there are two plants or only one. In crosses between dense and lax plants, in which the segregation is 3:1, wht uf wosibilities there are \(y\) chances

TABLE 21. Behavior of Fo Plants in F 3 . Series 13174 a Extra Early Windsor (Capitaticm) x Dale Gloria
\begin{tabular}{|c|c|c|c|c|c|c|c|c|c|c|c|}
\hline \multirow[b]{2}{*}{\[
\underset{\text { pedigree }}{\mathrm{F}_{2}}
\]} & \multirow[b]{2}{*}{\[
\underset{\text { density }}{\mathbf{F}_{2}}
\]} & \multirow[t]{2}{*}{Mean
density
of dense
\(\mathrm{F}_{3}\) plants} & \multirow[t]{2}{*}{Mean density F3 plants} & \multicolumn{2}{|l|}{Dense plants} & \multicolumn{2}{|l|}{Lax plants} & \multirow[b]{2}{*}{Total \(\mathrm{F}_{3}\) plants} & \multirow[b]{2}{*}{Deviation} & \multirow[t]{2}{*}{Prob able error} & \multirow[b]{2}{*}{\[
\frac{\text { Dev. }}{\overline{\text { P.E. }} .}
\]} \\
\hline & & & & Number obtained & Number calculated & Number obtained & Number calculated & & & & \\
\hline 34-3 & 1.31 & 1.29 & & 29 & & & & 29 & & & \\
\hline 31.8 & 1.40 & 1.28 & \(\cdots\) & 25 & \(\ldots\) & & \(\ldots\) & 25 & & & \\
\hline 34-11 & 1.55 & 1.36 & \(\cdots\) & 23 & & \(\cdots\) & & 22 & & & \\
\hline \(33-5\) & 1.55 & 1. 28 & & 46 & & & & 46 & & & \\
\hline 33-1.5 & 1.85 & 1.30 & & 46 & & & & 46 & & & \\
\hline 33-9 & 2.00 & 1.28 & & 39 & & & & 39 & & & \\
\hline 33-4 & 1.70 & 1.37 & \(3.09{ }^{-}\) & 59 & 64.5 & 27 & 21.5 & 86 & -+ 5.5| & \(\pm 2.70\) & 2.04 \\
\hline 33-2 & 1.74 & 1.57 & 3.77 & 56 & 63.7 & 29 & 21.2 & 85 & -+ 7.8 & \(\pm 2.69\) & 2.90 \\
\hline 34-4 & 1.86 & 1.57 & 3.69 & 31 & 32.2 & 12 & 10.7 & 43 & -+1.3 & \(\pm 1.92\) & 0.68 \\
\hline 34-6 & 1.95 & 1.54 & 3.31 & 23 & 24.0 & 9 & 8.0 & 32 & -+ 1.0 & \(\pm 1.65\) & 061 \\
\hline \(34-12\) & 1.95 & 1.69 & 3.33 & 23 & \(2 \pm 0\) & 9 & 80 & 32 & -+1.0 & \(\pm 1.65\) & 0.61 \\
\hline \(34-5\) & 2.00 & 1.39 & 3.32 & 18 & 17.2 & 5 & 5.7 & 23 & +-0.7 & \(\pm 1.40\) & 0.50 \\
\hline 33-6 & 2.00 & 1.54 & 3.59 & 49 & 51.7 & 20 & 17.2 & 69 & - 2.8 & \(\pm 2.43\) & 1.15 \\
\hline 33-7 & 2.00 & 1.60 & 3.34 & 45 & 49.0 & 19 & 16.0 & 64 & -+3.0 & \(\pm 2.32\) & 1.29 \\
\hline 33-3 & 2.00 & 1.60 & 3.12 & 43 & 50.2 & 22 & 16.7 & 67 & \(\underline{-+5.3}\) & \(\pm 2.39\) & 222 \\
\hline 33-13 & 205 & 1.60 & 3.80 & 47 & 487 & 18 & 16.2 & 65 & & \(\pm 2.36\) & 076 \\
\hline \(34-10\) & 2.11 & 1.69 & 3.48 & 23 & 24.0 & 9 & 8.0 & 32 & - 1.0 & \(\pm 1.65\) & 0.61 \\
\hline 33-17 & 2.20 & 1.57 & 3.53 & 49 & 52.5 & 22 & 17.5 & 70 & & \(\pm 2.44\) & 1.84 \\
\hline 33-11 & 2.21 & 1.65 & 3.50 & 51. & 570 & 2.5 & 190 & 76 & - + 6.0 . & \(\pm 25\) & 2.35 \\
\hline \multicolumn{3}{|l|}{Total heterozygous plants.....} & & 518 & 5584 & 226 & 186.0 & 744 & \(1-+40.01\) & \(\pm 796\) & 5.02 \\
\hline 33-10 & 3.30 & [... & 3.20 & . & .... & 22 & .... & 22 & & & \\
\hline \(34-1\) & 3.50 & & 3.35 & & .... & 34 & . . . & 34 & & & \\
\hline 33-1 & 3.71 & \(\ldots\) & 3 35. & & & 3.5 & & 35 & & & \\
\hline \(34-9\) & 3.93 & \(\ldots\) & 3.86 & & \(\ldots\) & 18 & \(\ldots\) & 18 & & & \\
\hline \(34-7\) & 3.96 & & 3.62 & & ... & 24 & & 24 & & & \\
\hline 33-16 & 405 & & 3.33 & \(\because\) & & 42 & & 42 & & & \\
\hline 33-12 & 4.42 & \(\cdots\) & 3.77 & & & 51 & & 51 & . & & \\
\hline \multicolumn{2}{|l|}{Total F3 plants} & & & & & & & 1,177 & & & \\
\hline
\end{tabular}
that a dense plant will grow next to a dense plant; 6 chances that it will grow next to a lax plant (or that a lax plant will grow next to a dense plant), and I chance that a lax plant will grow next to a lax plant. In other words, where entangling exists due to close planting, there will be six cases in which this condition will be detected and lax and dense plants separated, and nine cases in which it will be overlooked because the heads will all be dense and will show no visible difference. If this factor of entanglement is present whereby some dense plants are overlorked, theoretically there will be fewer dence plant- than are expected.

If the degree of experimental error introduced by this factor is calculated, it will be found that if, among 50 plants forming a segregating population, there are two or three cases in which a dense plant has beers interlaced with its neighbor and is not separated, the differences between recorded and theoretical ratios will be ahout as great as those shown in tables 20 and 21.

TABLE 22. Behayior of F2 Plants in F3. Series 13214a, Turiey (Vulgare) \(x\) Dale Gloria
\begin{tabular}{|c|c|c|c|c|c|c|c|c|c|c|c|}
\hline \multirow[b]{2}{*}{\begin{tabular}{l}
\(\mathrm{F}_{2}\) \\
pedigree
\end{tabular}} & \multirow[b]{2}{*}{\[
\underset{\text { density }}{\mathrm{F}_{2}}
\]} & \multirow[b]{2}{*}{Mean density of dense F3 plants} & \multirow[b]{2}{*}{Mean density of lax F3 plants} & \multicolumn{2}{|l|}{Dense plants} & \multicolumn{2}{|l|}{Lax plants} & \multirow[b]{2}{*}{Total \(F_{3}\) plants} & \multirow[b]{2}{*}{Deviation} & \multirow{2}{*}{Probable crror} & \multirow{2}{*}{\(\frac{\text { Dev. }}{\text { P. E. }}\)} \\
\hline & & & & Number obtained & Number calculated & Number obtained & Number calculated & & & & \\
\hline 38-8 & 1.72 & 1.46 & & 68 & & - & & 68 & & & \\
\hline 39-9 & 1.91 & 1.66 & & 45 & & & & 45 & & & \\
\hline 38-12 & 1.95 & 1.41 & & 97 & & & & 97 & & & .... \\
\hline 38-14 & 1.95 & 1.49 & & 85 & ... & & .... & 85 & & & \\
\hline 39-4 & 2.06 & 1. 66 & ... & 33 & . & & \(\ldots\) & 33 & & . . . & \\
\hline 38-11 & 2.16 & 1.51 & .... & 59 & . . . & . & ... & 59 & \(\ldots\) & & \\
\hline 38-9 & 2.33 & 1.61 & .... & 83 & \(\ldots\) & . & .... & 83 & . . . & & \\
\hline 38-13 & 2.39 & 1.58 & .... & 57 & & .. & & 57 & . . . & & \\
\hline 38-4 & 2.75 & 1.49 & & 44 & & & & 44 & & & \\
\hline 40-3 & 1.57 & 1.37 & 3.18 & 73 & 57.8 & 4 & 19.2 & 77 & +-15.2 & \(\pm 2.56\) & 594 \\
\hline 40-6 & 2.11 & 1.65 & 3.23 & 47 & 41.3 & 8 & 13.7 & 55 & +-5.7 & \(\pm 2.17\) & 2.63 \\
\hline 38-15 & 2.11 & 1.58 & 3.20 & 35 & 40.5 & 19 & 13.5 & 54 & -+ 5.5 & \(\pm 2.15\) & 2.56 \\
\hline 40-1 & 2.28 & 1.64 & 3.15 & 42 & 480 & 22 & 16.0 & 64 & -+6.0 & \(\pm 2.34\) & 2.56 \\
\hline 39-3 & 2.35 & 1.67 & 3.28 & 29 & 31.5 & 13 & 10.5 & 42 & -+2.5 & \(\pm 1.89\) & 1.32 \\
\hline 40-2 & 2.35 & 1.91 & 3.51 & 53 & 54.8 & 20 & 18.2 & 73 & \(-+1.8\) & \(\pm 2.50\) & 0.72 \\
\hline 39-2 & 2.53 & 1.76 & 3.44 & 52 & 51.7 & 17 & 17.2 & 69 & - + 0.2 & \(\pm 2.39\) & 008 \\
\hline 38-2 & 2.53 & 2.08 & 4.30 & 10 & 12.8 & \({ }^{7}\) & 4.2 & 17 & \(-+2.8\) & \(\pm 1.20\) & 233 \\
\hline \(40-7\) & 2.57 & 1.92 & 3.77 & 46 & 42.0 & 10 & 14.0 & 56 & \(+-4.0\) & \(\pm 2.19\) & 1.83 \\
\hline 38-10 & 2.61 & 1.96 & 3.51 & 30 & 30.0 & 10 & 10.0 & 40 & + 0.01 & \(\pm 1.85\) & \\
\hline 39-5 & 2.75 & 1.87 & 3.32 & 24 & 23.3 & 7 & 7.7 & 31 & +-0.7 & \(\pm 1.63\) & 0.43 \\
\hline \(40-8\)
\(38-5\) & 2.79
2.89 & 2.13
2.16 & 3.78
4.09 & 48 & \begin{tabular}{l}
585 \\
24 \\
\hline
\end{tabular} & 30
9 & 195
80 & 78
32 & -+10.5 & \(\pm \underline{1.58}\) & 4.07
0.61 \\
\hline 38-5 & 2.89
2.90 & 2.16
2.23 & 4.09
3.72 & 23
47 & 240
450 & 9
13 & 80
150 & 32
60 & -+1.0 & \(\pm 1.65\)
\(\pm 2.26\) & 0.88 \\
\hline 40-4 & 2.94 & 2.18 & 3.59 & 48 & 4.5 .8 & 13 & 15.2 & 61 & +-2.2 & \(\pm 2.28\) & 0.96 \\
\hline 50-10 & 3.00 & 2.20 & 3.83 & 51 & 488 & 14 & 162 & 65 & +-2.2 & \(\pm 2.36\) & 0.93 \\
\hline 39-6 & 306 & 2.28 & 3.89 & 73 & 713 & 22 & 23.7 & 95 & \(1+-1.7\) & \(\pm 285\) & 0.60 \\
\hline \multicolumn{3}{|l|}{Total beterozymous plants.} & & 731 & 7267 & 238 & 2422 & 969 & \(1+-4.21\) & \(\pm 9.09\) & 0.46 \\
\hline 38-3 & 406 & I .... & 3.94 & . & . . . & 37 & .... & 37 & & & \\
\hline 38-1 & 4.11 & .... & 457 & & & 12 & \(\ldots\) & 12 & & & \\
\hline 38-6 & 4.14 & & 3.75 & & & 41 & -... & 41 & & & \\
\hline \(39-1\) & 4.21 & & 3.41 & & & 32 & & 32 & & & \\
\hline 38-16 & 4.59 & & 3.78 & & & 58 & & 58 & & & \\
\hline 40-5 & 529 & & 3.64 & & & 91 & & 91 & & & \\
\hline \multicolumn{2}{|l|}{TotalF3plants} & & 1 & & 1 & & & | \(1 . \times 11\) & & & \\
\hline
\end{tabular}

In crosses I3If3a and I3If4a, density was the only visible differenthathe chancter; heme the speration of the entangling plant depended merely on that character. In series I3I72a and I32I4a, in which the experimental errors were practically as much on one side as on the other, the error due to entangling was reduced, respectively, by the introduction of the pubescent glume character through the Mealy parent, and by the color of chaff and the beardedness introduced by the Turkey parent. The introluction and comespuent nearesatom of these character- enahled the author to detect the presence of more than one plant, and through their separation the degree of experimental error in these two sets of crosses was greatly lowered.
lif comnection with tathe If 1 pase est 3 the reader perhaps noticed that there was a tendency on the part of some \(\mathrm{F}_{1}\) and \(\mathrm{F}_{2}\) generation freyumeles to be shifted somewhat toward the laxer clasess while whers tended to shift tonard the denser classes. since the dense parent , Dale Gloria, was the same in all seven cases, thee variations, if hereditary :u any extent, should he ascribed tu the influence uf lax parents which represent the variable factors.

The mean densities of parent and offormeng are represented graphically in figure So. The curves are arranged in the ascending order of the


Iitg. ©O. INFLLENCE OF DENSITY OF LAX PARENT ON DENSITY OF F1- AND F \(2^{-}\) generation Plants
den-itics of the lax parents. The straght lines fitted the the curves show a seneral rise; that is, with the increase of the average internode length of the lax parent, the average internode length of the \(\mathrm{F}_{1}\) and the \(\mathrm{F}_{2}\) increase more or less in the same proportion.

The slopes of the fitted lines for the \(\mathrm{F}_{1}\), the total \(\mathrm{F}_{2}\), and the dense \(\mathrm{F}_{2}\)
 spectively. Those for the lax parents and the lax \(F_{2}\) segregates show also a general rise but of a higher degree. It should be borne in mind that the higher the clas: values, the sreater is the tendency of the curve toward a higher inclination.

The curve repreanting the densities of the \(\mathrm{F}_{1}\) and the dense \(\mathrm{F}_{2}\) segregates follow each other very closely. The other curves also follow the same gencral conrec. Eidently the material repeesenting the cross with Mealy was somewhat denser, because both the \(F_{1}\) and the \(F_{2}\) curve show a similar rise at that point. The rise of the Mealy parent is not in the same proportion.
Aside from these differences, it should be noted that the values of \(\mathrm{P}_{1}\) and \(F\), are higher locause they repreant crope srown in different years and also because they were spaced more widely than the \(\mathrm{F}_{2}\) plants. Besides, the \(\mathrm{F}_{1}\) perlaph hows vieur due to heterozgens, which, together with increased food supply due to the wide distances between plants, tends to increase the size of the spike without increasing the number of spikelets, which in turn tends to increase the average internode length.

RELATION OF DENSITY OF I \({ }_{2}\) PLANTS TO THAT OF THEIR PROGENY
The comparison of the density of \(\mathrm{F}_{2}\) and of \(\mathrm{F}_{3}\) plants leads to the decision as to whether the variations in density, especially of the \(F_{2}\) heterozygous plants, are hereditary or represent mere fluctuations due to external conditions. To be sure, environmental conditions, as is pointed out in the first part of this paper, have a great influence on the degree of density. The plants used in this experiment were grown on a small area, and comequently the envirmmental factors hat practically as much uppromity to affect the demity of one plant as that of another.

In comparing the density of the \(\mathrm{F}_{2}\) plants with that of their progeny, series i32I4a may be taken as an example. Deductions based on this cross will apply as well as for the other crosses. The mean densities of
the \(F_{2}\) parent and the offspring, as shown in table 22, are represented graphically in figure 8i. Comparison of the density curves of the \(\mathrm{F}_{2}\) plants with the mean density curves of their progeny shows that, especially in the case of the heterozygous \(\mathrm{F}_{2}\) planis, there is a correlation between the density of the \(\mathrm{F}_{2}\) and that of their \(\mathrm{F}_{3}\) segregates. Since there is such a correlation between \(\mathrm{F}_{2}\) and \(\mathrm{F}_{3}\), these apparent fluctuations are


Fig. 81. Comparison of density of for With that of dense (homozigous OR HOMOZYGOUS AND HETEROZYGOUS) AND LAN PROGENY
hereditary variations. It goes without saying, then, that these \(F_{2}\) heterozygous plants are not genotypically identical with respect to compactness, and that, although all the \(\mathrm{F}_{2}\) heterozygous plants have the general formula \(C c\), they carry besides this factor a group of other factors which tend to increase or decrease compactness.

RELATION OF DENSITE OF DENSE AND LAN SEGREGATES OF HETEROZVGOUS
\[
\mathrm{F}_{2}, \mathrm{PLANTS}
\]

The presence of factors modifying the degree of density may be further demonstrated ble a comparton of the curven of the denne seeregatos
with those of the lax segregates. This test is made on the assumption that if a group of modifiers is introduced there will be as much chance for these to be transmitted to the dense segregates as for them to be transmitted to the lax forms of the progeny of heterozygous dense \(F_{z}\) plants; in other words, both dense and lax forms will receive the same duee uf modifiers. Accordingly, if a set uf modifiers shifts the mean of the dense segregates, say about five classes, toward a plus or a minus drectiom, the mean of the lax segregates should le shifted likewiee, to the extent of as many classes at least, and very likely more, toward the same direction.

On examining the density frequencies shown in tables 23,24 , and 25 , it will be noted that whenever the dense segregates of \(\mathrm{F}_{2}\) heterozygous plants are grouped toward the laxer classes the curve of the lax segresate correnanding te them tems aloo to arranse itedf in that direction, and vice versa. For example, in table 23 (series I3214a), of the progeny of \(40-3\) the dense forms are very dense, with a mean density (table 22) of 1.37 . The density of the lax form is 3.18 . Comparing this with the progeny of \(3^{\circ}-2\), the mean of whe dense segregate in 2 . of and of the lax 4.30 , it may be seen that whatever interfered with the compact-me:- of the dene plants of 3 s \(^{-2}\) a affected alon the density of the lax froms. Throughout tables 23. 24, and 25, in which the details of the frequency distributions are given, the same phenomenon may be observed. The rebation between the mean den-ity of the dense eegregates and that of the wresponding hax estesate of the F , progeny of each heterozysous
 is seen in these graphs that with the increase or decrease in the density of the dense plants, the density of the lax forms varies in the same direction. This is a direct evidence that besides the density factor there are alow modifier affectins the desper of demety within the deme and the lax classes.

GENERAL CONSIDERATION ON TIE FREQUENCY DISTRIBUTIONS OF COMPACT X LAX CROSSES
Each dense \(x\) lax cross has its own peculiar type of frequency distribution, either in the first or in later generations. The type of distribution seems to depend on the density of the parents concerned and on the


FIG. \&2. REIATION OF JIENSITY OF DENSE AND LAX SEGREGATES OF
HETEROZYGOUS FE FLANTS

Net ,if m, lifying factors introduced by them. In the discussion of table It ( pases \(\mathbf{s}^{5}\), it was bought nut that lax plants differ from one another in degree of density, and that the density of their progeny varies according to the density of the lax parent, the dense parent being the same. As In the que-tion of modifiers, the frequency-diotribution tables 23,24 , and 25 show the characteristics of the curves of each cross.

In series 1321 ta (table 23), althongh there is a gap letween the dense and lax curves, in a number of cases this gap is not so evident. In figurins wut the ratios the determination of the posible line of separation was a matter of judgment in some cases. Series 13Ij2a (table 25) shows a wider gap. To a very small degree the fewness of the plants may account for it. The plants of series \(13174 a\) (table 24) show a much wider gap and the presence of two definite curves is at once seen. In crower letween denee and lax forms, all gralation-between a distribution such as is shown in series I3174a, and a continuous skew curve, may be obtained.

Crosses made between semi-dense squareheads (often classified as clubs) and lax forms produce an uninterrupted curve which makes it practically impo-ible to eparate them into dense and semi-denee, and lax genetic, classes.

Kecrasling the mendelian clawification- of segregating line woh as those produced by the heterozygous \(\mathrm{F}_{2}\) plants of series 13214 a , it will not be out of place to make a few remarks. Some workers on this subject have taken a certain class of density as a dividing line between dense and lax forms, presumably based on the classes of least frequency of the \(\mathrm{F}_{\mathrm{a}}\) curve. While such a method may be more or less satisfactory in a cross similar to series 13154a (table 24), it is absolutely unjustifiable in genetic studies and unsuitable for the great majority of crosses in which dense or lax forms appear. In the first place, since \(F_{2}\) and \(F_{n}\) plants are grown in different years they do not necessarily show the same degree of density: This is shown in table 23. There is practically no heterozygous plant of the \(\mathrm{F}_{3}\) generation which produced a curve similar to that of the \(\mathrm{F}_{2}\). Thenretically about half of the \(\mathrm{F}_{3}\) curves should have approached the \(F_{2}\) curve. The reason for this failure lies in the fact that the \(\mathrm{F}_{2}\) plants were grown under more favorable environmental con-
SQuareiteadedness and Density in Wheat


SQuareheadedness and Density in Wheat

ditions than were the \(\mathrm{F}_{3}\). Now if, for example, density class 3.2 based on the \(F_{2}\) curve (table 23) is taken as the dividing line and is used for separating the dense and lax forms of the \(\mathrm{F}_{3}\), it will be noticed that a large number of lax forms of the \(\mathrm{F}_{3}\) will be included in the dense class.

It has been mentioned that each \(F_{3}\) curve has its peculiar mode of distribution. In tables 23 and 25 it is shown that the frequencies of some dense curves extend beyond the classes where the curves of some lax forms have begun. The peculiarity of each curve necessitates the taking of each curve and the separation of the dense from the lax forms at the class or the probable class of least frequency. If the density curve of segregating form are not prepared, more trustworthy reaults can be obtained he clansifing the forms bey ere than takine an arbiraty dividing line. This latter methed can le ued in case in which the variations are discontinuous.

In specices crosses, discontinuous curves are the exception rather than the rule, and separation of dense and lax forms into two genetic classes ly an arthitrary dividing line ha- no meaning becane such a separation cannot le made even ly the examination of the curver, even direcgarding the fact that some specific forms which carry the density factor may be laxer than most of the vulgare forms.

That density is a dominant character and is inherited in the 3 :I ratio
 Von Tchermak (1910), Nilsson-Ehle (igit), Mall (19i2), Parker (1914), and others. The three last-named have called attention to the fact that form- laner and dener than the parental types are ohtained also in compactum \(x\) vulgare crosses.

The use of the word squarchead for semi-dense forms by some authors accounts for the disagreement of their results. Von Rümker
 head, and vultgare forms in approximately the ratio I:2:1. No doubt the "syuarehead.," which proved to he heternzygous, were heternzygous clut) forme. Von kimber cite- anolher cave in which the \(F_{2}\) eegregated into dene and lax form-appromatins \(3: 1\), there beins no squarehead
 from dense forms.

Biffen's (1905) results, which are often cited as contrary to those obtained by others, seem rather to agree with Von Rümker's. Such differences as exist are due to methods and nomenclature. Biffen crossed squarehead's Manter with Ked King, the average internode length, being, respectively, 3.2 and 4.6 millimeters. The \(F_{1}\) hybrid ears averaged 4.8 millimeter: in internode length, and the Fo ranged from 3.2 to 5 millimeters. In this early work the frequencies are not given, but it appears that the distribution was of the normal, or skew, type. Using 4.6 (the mean of Red King, or the internode length of this lax parent) as the dividing line, Biffen fond that -8 flants were laxer than the lax prarent and 22 were intermediate between the dense and the lax parent, while there was no form denser than Squarehead's Master. From this result Biffen concluded that in this case laxness was dominant. In the first place, Squarehead's Master, with an internode length of 3.2, does not fall in the compactum class and the cross should not be included in that category. In the second place, the use of the density of the lax parent as the dividing line seems to be too arbitrary and unjustifiable from a genetic standpoint.

Neither does Biffen's Rivet \(\chi\) Polish cross fall in this category, since it does not belong to the sativum group. In crosses of this type specific ergregations occur which greatly complicate ratur. Hi: Dewn : Hedselog cross is of the semi-dense compactum \(x\) lax type. The photographs shown in the article cited, and also by Bateson (1909:23), show that the \(\mathrm{F}_{1}\) is intermediate dense and the \(\mathrm{F}_{2}\) evidently approximates the 3: I ratio, the heterozyon forms being slightly laxer than the dence parent.

Biffen's results, then, so far as the compactum \(x\) lax cross is concerned, may be regarded as being in accord with similar work done by others.

Since different types of dense and lax forms have been used by various investigators, their results with respect to degree of density have been different. Segregation into two distinct curves is obtained as the degree of density of the parents approaches the extremes of the dense and lax classes. As a rule, the closer the parents approach in degree of density, the more continuous the curves will be. This statement applies for crosses made within the sativum group. The existence of several factors producing varying degrees of density is evident.

\section*{TIE NATURE OF DENSITY FACTORS}

There que-tions of epecial interest ariee, regardine the nature of the den-ity factors producing varying degree of den-its in su-called pure lines. They are: (I) Is the density of the compactum type caused by hot one factor pair, and are the degreso of density produced by the presence of modifiers? (2) Do the forms varying in degree of density carry different density factor-allelommphic to one another, or (3) . Tre they caused by multiple factors?

A considerable amount of work has yet to be done to answer these questions definitely, an numernus senetic analyes are necessary. A few sugecotion: bacel on experimental roulti can be made. howeser, regarding the probable nature of these density factors.

The results of all the compactum \(\times\) lax crosses reported thus far show the presence of but one density factor; a \(15: 1\) ratio has not yet been obtained. But in crossing a compactum with a somewhat dense culgare, not inferpuently form- are whanch which are laxer than the cuthure and dener than the ompuctom parent. For the ake of simplicity in previous discussions, this occurrence was ascribed to the presence of modifiers. The writer believes that these modifiers are secondary factors representing varying degrees of density. The factorial combinations
 illustrated in the following manner:
\begin{tabular}{|c|c|c|c|c|c|c|c|c|c|c|c|}
\hline \(\mathrm{F}_{1}\) & \multicolumn{11}{|c|}{\begin{tabular}{l}
CC dd \(x\) cc DD \\
contpactum slightly dense zulgare
\end{tabular}} \\
\hline \(F_{1}\) & \multicolumn{11}{|c|}{\begin{tabular}{l}
CcDd \\
compactum slightly laxer than dense parent
\end{tabular}} \\
\hline \(\mathrm{F}_{2} \frac{1}{2}\) & \[
\begin{aligned}
& \text { CCDD } \\
& \text { CCDd }
\end{aligned}
\] & - & \[
\begin{aligned}
& \mathrm{CCdd} \\
& \mathrm{CcIDD}
\end{aligned}
\] & & \begin{tabular}{l}
Cend \\
CcDil
\end{tabular} & 1 & ccDD & & ccDd & & cocid \\
\hline \multicolumn{6}{|c|}{12 compactum} & & & & zulgare & & \\
\hline & \begin{tabular}{l}
denser \\
than \\
dense \\
parent
\end{tabular} & & \begin{tabular}{l}
approxi- \\
mately of \\
same doti- \\
sity as \\
dense \\
parent
\end{tabular} & & laxer than dense parent & & same as z'ulgare parent & & \begin{tabular}{l}
slightly \\
laxer \\
thatı \\
zulgare \\
parent
\end{tabular} & & \begin{tabular}{l}
much \\
laxer \\
than \\
vulgar \\
parent
\end{tabular} \\
\hline
\end{tabular}

In the preceding outline, \(C\) stands for the high degree of compactness introduced by the compactum parent, and \(D\) for the slight degree of compactness exhibited by the vulgare parent. This hypothesis accounts for the production of stable forms CCDD denser than the compactum parent and ccdd laxer than the vulgare parent.

In instances in which lax stable forms somewhat denser than the lax parent appear, the phenomenon may be explatnet acombling w the ame hypothe-is. If the above extreme dente ( \((1) /\) ) and lax codd forme taken as parents are crossed, stable forms denser than codd will be obtained according to the factorial combinations.

If there are different factors of density producing the compactum type, certain crosses between two compactum forms should give i in I6 or I in \(6 \neq\) lax forms. Let it be supposed that the compactum parents have the genotypic forms \(\mathrm{C}_{1} \mathrm{C}_{2} \mathrm{C}_{2} \mathrm{C}_{2}\) and \(\mathrm{C}_{1} C_{1} C_{2} C_{2}\) where there are two different \(C\) factors. With two factor differences, one plant out of sixteen in the \(\mathrm{F}_{2}\) should have the \(c_{1} c_{1} C_{2} C_{2}\) constitution, therefure being very lax. The writer knows of but one cross between two club forms, and from that cross, among \(\mathrm{I}_{3} 0 \mathrm{~F}_{2}\) plants no lax forms were observed, although considerable variations were found within, the dense classes. In this case both parents seemed to be homozygous for the same \(C\) factor; and the \(C\) factors contributed by both parents either were identical or belonged to the same allelomorphic series.

The author has evidence that the density factor of the compactum t: i" may letong to a multiple series. The black Pearded Spelt carries a density factor which may be isolated by crossing it with a lax vulgare form. About three-fourths to one-half of the \(\mathrm{F}_{2}\) vulgare forms are dense and semi-dense. In five such crosses the results have been similar. This shows that a \(C\) factor is carried by this particular spelt. If this factor is identical or forms an allelomorphic series with another \(C\) factor, the \(\mathrm{F}_{2}\) satioums segregating in a cross between this spelt and a club should be all dense. The fact that in such a cross lax satioum forms also are occasionally obtained, shows that in this case there were two distinct \(C\) factors involved.

Another \(C\) factor is carried by the dicoccum form known as Black Winter Emmer. When this form is crossed with a z'ulgare it throws
a certain proportion of distinctly dense sativum individuals. The \(F_{2}\) flants from eight such crosecs were examined and compact foms were recorded in every case. A= in the preceding instance, in the \(F\) : derived from two Black Winter Emmer \(x\) club crosses, consisting of abnent \(150 \mathrm{~F}_{2}\) individuals, soft lax vulgare types conld be counted, although most of the sutiabur type were dence. Here ascain it appears that, a in the preceding cros. two different non-allelomorphic mutiple \(C\) factors were introduced.

Summarizing the foregoing discussion, it is apparent that:
1. There are dem-ity factors cach producins shortening of the wachis impernode in different desrees. The en mas be preent in addition to the compactum factor.
2. There is as yet no sufficient evidence that some compactum factors may form an allelomorphic series. Neither is there any evidence that such series may not exist.
3. There exists in wheat more than one density factor belonging to multiple series. If such multiple density factors are present, wentually I5: I, 63 : I, and other ratios will be obtained from compactum \(x\) vulgare crosses.

FACTORS PRODUCING SQUAREIIEADEDNESS AS COMPRISING ONE OF THE GROUP OF FACTORS MODIFYIING DEGREE OF DENSITY
A modifier, as generally held, may be a factor affecting a particular character quantitatively, the presence of which is detected from the degree of modification of the character which it modifies. In a broad sense, however, any factor or character whose presence affects or even
 sarded as a modifier.

The irregularities among the \(F_{a}\) density curves of dense \(\underset{\sim}{\text { lax }}\) crosses shown in this study were ascribed to the presence of modifiers, ia the senetic analyes applied the the crones showed conclunively the presence of but one density factor. Proofs that these variations were Berelitary, and not caused be envirommental conditions, were alow riven.

The nature of one of these modifiers, the character for squareheadedme.. mat here le com-ldered. In (rwne in which a non-upuarehead was involved, it was noticed that there was an independent segregation of
density and -quareheadednes, and that whenever the latter character wan present, whether the plant was dense or lax, there was, as a rule, a reduction of the average internode length.

Density notes derived from series i3sza are summarized in table 2th. Thi, cross, it will be recalled, is between Mealy, a lax mm-apuarehead,

TABLE 26. Mean Density of \(\mathrm{F}_{3}\) Lax Plants Classified According to Proportion of Squareheads and Vulgare Forms. Series I3If 20

and Dale Gloria, a dense squarehead. In this cross both density and squarehearledness segregated independently. The table shows the density of the lax forms of the \(\mathrm{F}_{3}\)-generation lines arranged in five classes, accosding to the observed propurtions of squareheads to nom-splatreheads. In the first class are included the \(\mathrm{F}_{3}\) plants which consisted of squareheads only; the second class includes the pedieree that prombed more squareheads than vulgare forms; the other three classes represent progenies consisting of an equal number of these two forms, of an excess of vulgare, and of only vulgare, respectively.

By averaging the degree of density of each group it is found that the average internode length increase invernely with the proportion of squareheads. The average internode length of the squareheads was \(3 .(1)=0.04\), while that of the plants producine an excen of spuareheats was \(3.28 \pm 0.07\). The densities of the class yielding more vulgare than squareheads and that producing only zulgare were \(3.72 \pm 0.11\) and \(3.7 \mathrm{I} \pm 0.05\), respectively. The difference in the average internode length of the extreme classes was \(0.65 \pm 0.06\). Comparison in terms of the average of the means of these two classes, shows this difference be-
tween the density of fure - puareheads and that of pure erflewt to be equal to about 20 per cent of their means, which is rather remarkable.

Were the lax plants of the \(\mathrm{F}_{3}\) generation all or practically all squareheads, one would expect the elimmation of variations as great as those found in the crosses between vulgare and dense squareheads. Series I3174a (table 24, page 850), which represents a cross between a squarehead and a dense -quatehead, show: the alsonce of thifting of the curves back and forth, so evident in tables 23 and 25, which represent dense squarehead \(x\) vulgare crosses.

Although the variations introduced by the presence of both vulgare and squarehead forms is eliminated in cross I3Ifta (table 24), other montifer mu-t le present becane the cume still show imheritable variations. The nature of these renaining modifiers is not as yet known.

It is not difficult to explain how squareheadedness increases density. Ihn-whosical studien show that suareheadednes. which is brought about by the shortening of the terminal intemodes, is due to contact and pressure caused by the differential rate of growth of parts of the plant surmonding the fuke during its earlier periods of srowth. If, during the process, the terminal internodes fail to attain their normal size, that part of the head will be denser than it would have been if the plants were allowed to grow normally. This shortening of internodes of the lerminal part of the head is the direct cause of the decrease of the average internode length of the entire head.

RELATION OF SQUAREIJEADEDNESS TO DENSITY IN F \({ }_{2}\)-GENERATION PLANTS
The modes of inheritance of squareheadedness and of density have been discussed separately herein. It is necessary now to illustrate the relation between these two characters in crosses in which both of these characters have been introduced. Series 131ヶ2a, \(13214 \mathrm{a}, 1337 \mathrm{a}\), and 13153a (tables \(27,28,29\), and 30, respectively), which have already been considered, will be used again as examples, because each illustrates a different mode of inheritance.

Before examining the behavior of the \(F_{2}\) plants, the distribution of the parent plants with respect to density and squareheadedness may be recom-idered. Thi di-tribution, althnigh hated on few mumbers. is hown


Squareheadedness


Fig. 83. distribution of parental plants with respect to DENSITY AND SQUAREHEADEDNESS
the varietics used in making the crosses; they are the progeny of each of the plants used as parents.

Dale Gloria, it is seen, is distributed within a very narrow range with respect to density, hut within a wide range with repeet to spuareheadedness. This latter range does not represent the genotypic range of the variety, if one may be allowed to use that expression. The length of the rachis itself is about 2 to 3 centimeters. If the terminal spikelets are ster-
ile, a, often happens in dense clubs and lese frequently in lax forms, this sterility and rudimentary condition will keep the terminal rachis internodes from developing further while the other internodes continue their growth. A head of this type will have a high coefficient of squareheadedness. If, on the contrary, the terminal Horets develop vigorously, the increase in the size of the grain will tend to stretch the internodes somewhat. With a vigorous growth of the grain is associated a relative grow of of the adjacent internodes, and with as short a head as that of the cluh in question it doe not take much increase in internode length to lower the squarehededness of some individuals down to classes 1.00 to I. 25.

The distribution of New soules is entirely different from those of the dher lax plants used in the crosses. It is somewhat more oward the demer classes and falls distinctly within the squarehead clases.

Turhinh Amber is a coulnore, but its average internode length is greater than that of any other of the forms represented.

TABLE 27. Relation of Squareheadedness to Density in \(\mathrm{F}_{2}\) Segregates. Series 13172a, Mealy x Dale Gloria
(Mean dense plants, \(\mathrm{D}=2.01, \mathrm{Sq} .=1.34\); mean lax plants, \(\mathrm{D}=3.95, \mathrm{Sq} .=1.23\) )


Mealy has practically the same distribution as Turkish Amber but is somewhat less lax.

Turkey is the least lax in the vulgare group and is located very much toward the non-squarehead classes.

The mode of inheritance which might generally be expected in compactum \(x\) eulgare crosses is that of the Mealy \(x\) Dale Gloria series (table 27), which shows a wide range of distribution with respect to squareheadedness. Thenretically, since Dale filoria carres the factor- re. sponsible for squareheadedness and Mealy does mot, both density and squareheadednes would be expected to segrepate independently, which happens in this case. Using, for practical purposes, class I. 30 or I. 35 as a dividing line, it is seen that there are about as many individuals on the side of squareheadedness as on the side of non-squareheads. By using as a guide the distribution of the Mealy parent shown in figure \(\mathrm{s}_{3}\), it becomes very evident that in this cross the character of squareheadedness has been introduced among the non-spuareheads. The mean salue of squareheadedness of the lax parent is I.If and that of the lax plants of the cross is 1.23 , showing an increase of 0.09 . While out of 15 Mealy parental plants (figure \(\mathrm{S}_{3}\) t there wa- \(1,01 t\) one individual in the squarehead classes in the \(\mathrm{F}_{2}\) generation, out of a total of 20 plants there were about four or five times more individuals in these classes. In table 26 , which represents the same cross, it has been shown that some of these lax squareheads remained stable.

The distribution when Turkey was the lax parent (table 28) was strikingly different from that in the case of Mealy. In the Turkey \(x\) Dale Gloria cross, no lax squareheads appeared.

On comparing the dense classes of tables 27 and 28 with respect to their squareheadedness distribution, it is seen that the entire distribution has been shifted to the left in the latter table. The mean squareheadcdness of Dale Gloria is I .28 ; that of the dense plants of the cross is I.I7, showing a shifting of O.II toward the non-squarehead classes. While the squareheadedness of the dense form in affected by that of the lax forms, the difference in the coefficients of the lax parent and the
 are in turn influenced to some extent by the squareheadedness of the dense parent.

TABLE 28. Relation of Squareheadedness to Density in \(F_{2}\) Secregates. Series 13214a, Turkey x Dale Gloria
(Mean dense plants, \(\mathrm{D}=2.18 . \mathrm{Sq}=1.17\); mean lax plants, \(\mathrm{D}=4.23, \mathrm{Sq}=1.00\) )


In the crose in which Turki-h Amber wat used (table 29), the - , uare
 a- that of the lax pratent itself, the values of spuareheadednes of the lax parent and the lax segregates being I.O4 and I.O7, respectively: This - light hiftins toward the non syuarehead cla-ses may well le disegard\&.1. Unlike the combliom in the preceding catses, the dense fomm- showed no visible effect of the las parent. The mean spuareheadednes of the dense segregates, instead of being less, was slightly greater than that of the dense parent and practically the same as in series I3I72a.

In the fourth cross (table 30) the lax parent was New Soules, a dissinct squarehead. The lax segregates of this cross were all squareheads, and with regard to density they occupied the same position as the lax squarehead parent (figure 83 ).

Incidentally it should be noted from this table that, while the range

TABLE 29. Relation of SQuareheadedness to Density in \(\mathrm{F}_{2}\) Segregates. Series 1337a, Turkish Amber x Dale Gloria (Mean dense plants, \(\mathrm{D}=2.23, \mathrm{Sq} .=1.35\); mean lax plants, \(\mathrm{D}=4.30, \mathrm{Sq} .=1.07\) )

of density of the dense segregates of other crosses extends as far down as class 3.1 , that of this cross (table 30 ) is confined within classes 1.3 to 2 . I with but one individual in class 2.5. This difference is accounted for by the higher degree of density of New Soules (figure 83) as compared with those of the other lax forms.

Recapitulating the points of interest brought out in the discuss:on of these four tables, the following general deductions can be made:
I. The factors of density and those of squareheadedness are totally independent of each other. They segregate and recombine independently.
2. The wheat varieties of the satirum group have different degrees of density and of squareheadedness. Even among the oulgare plants this latter form of compactness, as measured by its coefficient, varies. In

\section*{fable sin Relation of Surareheadedness to Density in Fe Segregates Series 13173 a, New Soules x Dale Gloria \\ (Mean dense plants, \(\mathrm{D}=1.64, \mathrm{Sq}=1.43\); mean lax plants, \(\mathrm{D}=3.64, \mathrm{Sq} .=1.59\) )}

 lieats nor all dense forms produce a frepuency distribution of squareheadednes or of density of the same type and within the same ranse. Whether the range will remain apposimately in the same location, or will hift one way or the other, in determined to a great extent be the d.eree of squareheadedne or of densit! of the different forms of it.her or squareheads, as the cane may be, which are involved in the cross.
3. There are exceptions to the general rule stated above. In series 153, a there was a visible segregation of density but not of spuareheadednes. In this case there was evidently interference by another factor.

Sil-om-Ehle (19II) considers the inheritance of squareheadedness and give a factorial explanation to account for the apparent proportion in which it appeared in wome of his crosees. But, while Nilson-Ehle une 1 the term squarehed for lax forms which have a comparatively shorter average intemode lemeth than the culdure forms-that term is applied in the present paper to fomm: showine a relative density of the
middle and the upper third of the spike of about I. 33 or over, irrespec-
 son-Ehle's results and hypothesis with those from this study.

\section*{RELATION OF LENGTH OF RACHIS TO DENSITY IN HYBRID PLANTS}

Length of rachis is dependent on two factors, namely, the number of internodes and their length. If the number of internoder in at and. tion is more or less constant, as has been the case in all the crosses betheen Dale Ciloria and other forms considered hevein, the lemeth, if the rachis is directly proportional to the average internode length. This is so obvious that it needs no illustration.

If both factors are made variable by the selection of parents which vary both in number and in length of internodes, then there is no correlation between length and density: te an illustration a cron may'e cited which was made by the writer for this purpose. This cross was Silver Club x Aegilops arata. Silver C'luh, I Flate I. XIII, upper, 子: a clute wheat from four to five centimeters lons, with about ecentecn it twenty internodes. The Aegilops (Plate LXVII, upper, I) also was short, like the clitb wheat, but had only six internodes of an average lengeth of about six to seven millimeters. The hasal internode wewe the shorter, their spikelets being rudimentary.

Unlike the parent, the \(\mathrm{F}_{1}\) plants all hat hum head. reambline \(13: 0\) spelt wheat. Three plants obtained in the \(\mathrm{F}_{2}\) also were lax. The point of interest in this cross was that the \(F_{1}\) and \(F_{2}\) plants did not inherit length of rachis from their parents. Wut momber and kngh wifneme de. The \(F_{1}\) heads usually had from twelve to fourteen internodes from sixto seven millimeters in length. The three \(\mathrm{F}_{2}\) plants showed some variation in length. In the \(\mathrm{F}_{1}\) 'plants, both character lemes intermediate, the heads were necessarily much longer than in either parent. It would be expected, if sufficient \(F_{z}\) plants were eltamathe, that the phant- wend segregate with respect to both character intw bort head- deme an! o a that is, with many and with few internotes, and alo. comparation? 1 :n heads dense and lax, with possible intermediate forms.

In this connection it may be pointed out that what has been called vigor due to heterosis in wheat is often the appearance of musually long heads in \(F_{1}\) or later generations in crosses with certain emmers.

The ee are an a rule, smewhat dense and bear some thirty internodes to the -pike. Hence the question is rather one of number of internodes and internode length. The plants that combine the internode length of the culgare parent with the number of internodes of the emmer must necessarily be unusually long.

In actual practice, in a cross such as the above a considerable number of suthetic spelt, appear. These spelts have the peculiarty, as is shown later, of producing internoles lonser than those of the lax parent. The appearance of this new type helps to increase the proportion of unusually long heads.

RELATION OF LENGTH OF CUI,N TO RACHIS LENGTH AND DENSITV
In a pure line of wheat there is practically no correlation between culm lensth and demity, but there is a comelation between culne length and leneth of rachis. The plant that prodnce a short culm due to unfatorable envirommental conditions maturally produce a small head; but such a head as a rule has fewer rachis internodes than the mean of the line, and therefore, although the undeveloped head is short, its den-ity has mot been affected to any extent lecanse the number of rachis internodes has decreased more or less proportionately.

The writer's studies of the relation of culm length to density were mate on ceries izstata, theause this line produced practically no square-heal-. Squarcheading, it has leen -hown, unless it is due to faworable wrom th comditions, has a tendency to shonten the average rachis internode lemeth. There leems no equareheads in the material ued, that factor was eliminated.

The question of the relation of culm length to other characters of the home in of intered from both the commonic and the genctic viewpent. Fowathe of the many phase to which thi problem of density has led, it was not possible in this investigation to study the relation of culm whatere as inten-ively and extensively as the subject deserves. Sufficient data have been ohtained, however, to give an idea of the general behavior of this character.
bue (1) the great variability of culm length, the preliminary studies were makle with a mumber of progenies of \(F\), plants each of which had yielded on an average about 60 individuals. The frequencies of the

thete character-, culm length, rachis lengh, and rachis internode lensth, d.e hown in tathe 31 , tosether wath their means. The \(F\) lines inciuded in this tathe are arranged according to the order in which they were planted- \(3^{8-3}, 3^{8-4}, 3^{8-5}\), and so on, representing successive rows of flantings. In inspection of the table shows that these three characters - lengeth of culm, lengeth of rachio, and aterage internode length-are more or less closely correlated. As a rule the lines with long culms profucul lax plants and thome with hort culm- velded dense plant- It is




Fig. St. ajerage culm, rachis, and internode length of some f3
fanilies of Series 13214a
The curves show the relation of culm length to rachis and internode length. The order of the families corresponds to the order in which they were planted
when the mean of one of the characters decreases or increases, the others vary as a rule in the same direction.

There being no inherited variation in number of internodes (which averages about 20 to the spike) in the \(\mathrm{F}_{3}\) lines, the curves of rachis length and average internode length follow each other very closely. The curve of the culm length, although in a general way varying with density, shows certain irregularities, especially in the case of lines 38-6 and 40-4. The former is a homozygous lax type and the latter is heterozygous; yet the mean culm length of the former is considerably less than that of the latter, although theoretically it should have been greater. The possibility of the effect of environmental conditions being out of the question, it appears that there is a segregation of culm length independent of the dwarfing caused by the presence of the \(C\) factor. This statement is made as a suggestion only, since at the present time no definite explanation can be given.

A fact which seems rather interesting is that the \(C\) factor does not shorten the culm length in the same proportion as it shortens the rachis length. The averages of the rachis and culm lengths of the homozygous lax (cc) plants were about 7.7 centimeters and 89 centimeters, respectively. The presence of the double dose of the \(C\) factor (in \(38-4,3^{8-8}\), and \(38-9\) ) shortened the rachis length to an average of 3.2 centimeters and the culm length to about 74 centimeters, a shortening of 58 and 17 per cent, respectively, from the general average. In other words, while the rachis length was shortened by the \(C\) factor by more than one-half, the culm length was shortened but one-sixth.

Studies of the characters of dense and lax segregates have led the author to believe that the \(C\) factor is a dwarfing factor, shortening, besides the culm length, the rachis length, and the rachis intemode length, a number of other characters such as length of glumes, length of kernels, length of awns, and length of culm internodes. These two last-named characters have not been studied carefully by the author. Sapehin (1gnt, and his collaborators, who - tudied the ermedation between den-ity and culm internode length, claim that there is a significant positive correlation between these two characters. From the present studies it seems apparent that the shortening of the culm as a result of the pres-
ence of the factor for density, is due not to the reduction in number of cum internoles, but to the reduction in length of the culm internodes, the number of these internodes remaining more or less constant. In this repect the phenomenom oi the shortening of the culm is similar to that of the shortening of the rachis. It has been demonstrated by variou, workers that in maize aloo dwarfing cathe the shortening of the internodes of the stalk without necessarily affecting their number.

In a general statement such as is made here regarding the presence of a correlation between density of the head and hortness of the culm, it is not intended to convey the idea that dense plants or varieties are all tw :cort, and lax plants tall. The cardinal fuinth brought out are (I) that when the factor of density or its absence has been introduced in a proseny through hatridization, porided there are no interfering factors, the dense plants will be more likely to have short culms than the laxer plants: and \((2)\) that this bortening of the plant is caused, not by the reduction in number of culm internodes, but by the reduction in their length. It should be borne in mind, however, that these characters are affected by environment. From a genetic viewpoint the exhibiture uf a quantitative character in an individual plant is of little value especially if this is affected by enviromment. The comparative height of a plant is determined by the behavior of its progeny.

CORRELATION BETWEEN AVERAGE INTERNODE LENGTH AND LENGTH OF STFRIIE GLUMES

One of the proofs that the density factor is a dwarfing factor is found in the high degree of corrclation existing between the average internode lengeth and the leneth of the sterile glume- The material for the study of this correlation consisted of the spelt plants of series \(13255 \mathrm{a}-\mathrm{I} 5\), representing a cross between Dale Gloria and White Spelt. This \(\mathrm{F}_{3}\) line segregated into dense and lax sativums and spelts. In the data here only the spelts are represented. The spelts were selected primarily because the glumes could be readily removed from the spikelets, as they break oft uniformly at the base of the glume at a definite region just below the heel. With zulgare forms the taking of measurements is somewhat more laborions. The measurements of the glumes recorded here rep-
reant the average of the length of（opermens sterile glume on the sann spik：let at a distance from the bawe of the spike of about one－thed the length of the rachis．This precaution was taken because the glumes shorten as they approach the distal or the basal part of the head．

The correlation between average internode length and glume lengtla is represented in table 32．The correlation coefficient here is \(0.838 \pm\)

TABLE 32．Correlation between Density and Lengih of Sterile Glumes （Series \(13255 a-15\) ，Dale Gloria x White Spelt；only the spelts measured）

Length of glumes（in millimeters）
\begin{tabular}{|c|c|c|c|c|c|c|c|c|c|}
\hline & 6.5 & 7.0 & 7.5 & 8.0 & 8.5 & 9.0 & 9.5 & 10.0 & 10.5 \\
\hline \(\bigcirc 1.5\) & 1 & 1 & 1 & & & & & & \\
\hline 號 2.0 & & 3 & & & & & & & \\
\hline －2 2.5 & & 3 & 13 & & & & & & \\
\hline \％\％ 3.0 & & 1 & 1 & 3 & 2 & & & & \\
\hline ㅇ．． 3.5 & & & & & 1 & & & & \\
\hline 离䨌 4.0 & & & & & 2 & & & & \\
\hline \％ 4.5 & & & & 1 & 3 & 1 & 1 & 2 & \\
\hline 운의 5.0 & & & & & & 1 & & 1 & 1 \\
\hline \％ 5.5 & & － & & & & & 1 & 1 & \\
\hline 4 6.0 & & & & & 1 & 1 & & & 1 \\
\hline
\end{tabular}
0.039 ，which shows significantly that in this particular crow the factur decreasing rachis internode length is the one cansmg the shortening of glumes．The relation of density th glume length may be readily seen in heads 1 ， 7 ．and \(S\) in Plate LXIII（upper），which show the grade of density and consequently of glume length．
CORRELATION BETWEEN AVERAGE INTERNODE LENGTH AND LENGTH OF KERNELS
The same degree of correlation exists between density and length of kernels as between density and glume length．The measurements of the length of the kernel as here recorded represent the average leneth of the first and second kernels develned om the baval florets．（of there two） kernels the first was very often longer than the second．In case：in whet either the first or the second hasal thoret hat mot ponduced seed，the thend seed was not measured in its steat hecaure the thied seed is alwa：＇ik ly
to be -maller. In such cases a different -pikelet was chosen, the samples being taken always at a distance from the base of the head of about one-third the length of the rachis.

Correlating these two characters as shown in table 33, a correlation
TABLE 33. Correlation between Density and Length of Kernels (Series 13255a-15, Dale Gloria \(x\) White Spelt; only the spelts measured)

Length of kernels (in millimeters)

coefficient of \(0.747 \pm 0.024\) is obtained. Compared in terms of the respective probable errors, this is as high as or slightly higher than the correlation between density and glume length.

The width of the kernels was not affected by the length of the kernels or by density. This density factor does not seem to produce diminutiveness but to shorten along one axis only. The width of the kernels, whether taken laterally ur dur-i uentrally, remaining practically the same, the shape of the kernels assumes a spheroid form among the dense segregates and a long spindle form among the lax plants.

Since density is correlated with length of glame and length of kernel, it is obvious that in this material there is also a direct correlation between length of glume and length of kernel.
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    RELATION OF DENSITY OF RACHIS TO DENSITY OF RACHILLA
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A rachilla is the rachis of a spikelet, and branches out from the main rachis. A spikelet may have three or more rachilla internodes; the basal one is very short, but toward the terminal part of the spikelet the internodes elongate and then shorten again.

It is practically impossible to measure the average rachilla internode length. For comparative purposes, however, the relation of density to average rachilla internode length can lo determine indirectly by moting the extent of the protrusion of the fertile glumes of the florets. The relative distance between the fertile glumes of the first and the third floret on different wheat heads indicates their relative rachilla internode length, as is illustrated in figure 85 .


Fig. 85. spifelets of spelts of varying degrees of density, showing relation of
 and relative rachilla internode length as determined by mistance between THE TIPS OF THE FERTILE GLUMES OF THE FIRST AND THIRD FLORE1S \(\left(z, z^{\prime}\right.\), AND \(\left.z^{\prime \prime}\right)\)

To bring out the correlation between rachis and rachilla internode length it is necessary to find a population comprising wheat plants of the same species which are segregating into dense and lax forms.

Dense and lax spelt plants of series 13255 a -1 5 are reprersented, respectively, in \(A\) and \(B\) of figure 85 . The illustration shows that the laxer was the form, the more did the florets protrude above the two sterile glumes. In figure \(85, C\), is represented the appearance of a synthetic spelt of un-
usual length derived from a durum x vulgare cross. It shows a further increase in rachis internole kensth, whether with a relative increase in glume length.

These observations, which unfortunately cannot be presented in the usual form of a correlation table, indicate that \(C\), the factor for comfactues, women- alo the lensth of the rachillae or that of their internodes.

THE FACTOR IOR SPELTING ACTING AS A MODIFIER F゙OR THE DENSITY FACTOR
Density, like squareheadedness, is affected to a large extent by the presence of the spelt factor. In series 13255a, which represents a cross
 is but one spelt factor. Soplts and sationth.s segregate in this cross in the simple monohybrid ratio of 3 spelts of all grades (heads 6,7 , and 8) to I sativum (which includes compactum theads 9 and 10], squareheads thead II], and zullgare [head 12]). In a cross in which one of the parents in a spolt, the inheritance of density cannot be studied if all the plant- are clansified accordins to density alone. for, as will be seen, in the presence of the factor for spelting the factor for density does not produce compactness in the same degree as it does in the absence of the spelt factor. Therefore, in determining the mode of inheritance in such cases, it is necessary to take into consideration both the degree of spelting and the den-ity, and in interpeting the data the density curves of the spelt and those of the sativums should be examined separately.

The difference in density of the spelt and the satioum form may be loot illu-trated loy the density curve of the progeny of the two \(F\). flants, one of which was homozygous dense and the other was homozygous lax, but both of which segregated into spelts and sativums. The relative density of spelts and sativums is shown in table 34. The plants of 13255a-26 are segregates from the Dale Gloria x White Spelt cross. They are all homozygous dense, but are derived from \(F_{2}\) plants heterozygous for the spelting character. If \(S\) stands for the spelting factor and \(C\) for compactness, the \(F_{3}\) progeny of line \(13255 a-28\) consists of SScc, Sscc, and sscc individuals, while \(13255 a-26\) consists of SSCC,

for Sprlitiv,


SsCC, and ssCC plants. The mean internode length of the lax spelts, as shown in table 34, was 4.22 millimeters; that of the sativums was 2.56 millimeters, showing a difference of I .46 . Of the plants homozygous for the density factor, the dense spelts had a mean internode length of 2.00 millimeters and the dense sativums one of 1.57, showing a difference of 0.43 millimeter.

From these results it is seen that, although the internode length of the spelt can bee slortened ly the introduction of the ( factor, the preence of the \(S\) factor tends to interfere with the effect which a known \(C\) factor would otherwise produce.

The relation of the \(S\) factor to density is the same as the relation of this same factor to squareheadedness. It has already been shown that the presence of factors for squareheading have a very slight effect on the heads that carry the factor for spelting.

\section*{TIE SYNTHETIC PRODUCTION OL゙ TRITICUM COMPACTUM}

Triticum compactum, as has already been shown, is but a form of \(T r\), sativum, which carries an additional factor or factors for compactness. The results of crosses between different species show that this same factor can be carried as well by any of the other species of Triticum. In fact, there will be found in commercial strains species that carry density factors, though not necessarily the same factor. Many forms of durum, and especially of dicoccum, carry a density factor. When these are crossed with a lax zulgare form, compact forms invariably appear in varying proportions, and, depending on the genotypic forms of the plant, some of these compact forms breed true while others segregate.

In order to produce compact forms, it is not necessary that one of the parents should be a dense form. If the plant is carrying an inhibiting factor besides the factor for density, it may appear lax although it has the potentiality of producing dense forms. Thus, Black Bearded Spelt, which has been used by the author, is perfectly lax, but when crossed with vulgare it produces lax spelts in the \(\mathrm{F}_{1}\), and in the \(\mathrm{F}_{2}\) an apprecialle mumber of perfectly dense fomms. The White Winter Sielt, on the other hand, produces me compact forms. The Black Bearded Spelt, then, carries a factor for compactness and also an inhibiting factor. Certain
forms of turgidum, durum, and even polonicum also have the ability to produce dense types in the \(F_{2}\).

The writer has had no experience with vulgare crosses which have produced compact forms in the \(\mathrm{F}_{2}\); and unless the inhibiting factor is closely linked with the factors producing a certain specific form, it sempossible that certain vulgare forms will be met with which, although carrying the density factor, will be lax because of the presence of an inhibiting factor.

The mode of inheritance of density in some species crosses is very complex. because new specitic forms ariee in such ctanser, cach w which is affected in a peculiar way by the density factor. Besides factors for inhibition, many modifiers also may be involved. Often irregularizies are produced by the failure of development or maturation of some forms which seem to carry combinations of growth factors that restrict growth or cause various anomalies. Before being able to analyze from a factorial point of view the inheritance of density in such crosses, it is necessary to know in what proportions the various specific forms segregate.

There are two cases which suggest that compact or semi-compact vulgare forms have been obtained through crossing two vulgare forms. De Vilmorin (1913) cites two instances in which lax forms produced dwarf forms. The dwarf plant, a photograph of which accompanies the text, appears to be a compact form much denser than the plant from which it mutated. It may also be possible that these cases were natural crosses with a compactum pollen, since segregation of other characters occurred also. Another case is that mentioned by Neethling (1918), in which tall oulgare forms croned among themetves viedted dwarf form. The latter behaved as recessives. The statement is made that the dwarf plants had short ears, but nothing is said about density. If these were compactum forms, the fact that these dwarf forms appeared in a proportion somewhat less than 25 per cent tends to lend support to the possibility that the parent which carried the density factor carrime alon a factor inhibiting the production of dense forms. Until more is learned about the behavior of the \(F_{2}\) in the \(F_{3}\), no definite explanation can be given to account for its mode of inheritance.

In Plate LXVII (lower) are shown a number of dense forms of different wheat species, most of these being synthetically produced in interspecific crosses. Some show both compactness and squareheadedness; uthers show one of these characters in the absence of the other. Kürnicke (1885) has observed dense and squarehead varieties in other wheat species. He gives the following botanical varieties: Tritioum durum Desf. var. compactum Sér., Tr. polonicum L. var. compactum Link., Tr. polonicum L. var. quadratum Sér., and Tr. turgidum L. var. quadratum Sér.

Density, or the excessive shortening of the rachis internodes, is not contined to the genus Triticum. Dense forms are common buth in larley and in rye.

The question of the origin of Tr . compactum becomes simplified if it is recalled that this form may be produced when a sutituan form is cronsed with another wheat species, and also that natural eroseing vccurs not infrequently. Such being the case, one would expect Tr . ampuctun to lee practically a. old a form as any of the other species, and, so far as archaeological evidences go, cultivation of Tr. compactum has heen traced as far back as the stone Ase. Buschan (1895) state- that this compact form has heen found in the remains of caves and lake dwellings and among other prehistoric relics in regions extending from Egyt to central Europe and to Sweden. According to Unger (i860), the culture of wheat has been traced back to the year 3623 B. C., and hence its origin must be older still.

If interspecific crosses between vulgare and other forms are able to produce compact forms, it vem- that the first erigin of Tr. compuctum -hould have followed that of Tr. and!are. Undoubtedly Tr. compactum has reappeared many times in the same manner, for the afrearance of this form in interspecific crosses is rather common. Tr. compactum may le a mutational form of oulgore, although there is no defendable widence regarding this posibility for zulyare wheats. There is a parallel cample in the ca*e of the rye known an "Heinrich-Roggen" (Hillmann, 1910:579). This is a very compact form of rye which is said to have appeared in 1880 as a mutation on a single ear.

\section*{SUMMARY}

The density studies reported herein were made primarily on the progeny of a number of crosses in which the dense parent was Dale Gloria (Triticum compactum), with an average internode length of about 1.4I millimeters.

Density was found to be dominant over laxness. 'The ratios obtained approached \(3: 1\). The heterozygous forms were somewhat laxer than the homozygous dense forms, but hem nean- intermediate tetnetm :he dense and the lax parents. The \(\mathrm{F}_{2}\) curves were bi-modal and discontinuous.

The \(\mathrm{F}_{3}\) plants showed various degrees of density within the dense and the lax classes. Proofs are given in the text showing that these variations are hereditary and are the result of the segregation of modifiers or of additional density factors capable of producing density only within short ranges.

Experimental evidence is cited suggesting that different density factors form allelomorphic series, and other evidence that they belong to multiple series.

Squareheadedness and density were found to represent two different characters. Hybrid progenies showed all types and grades of combinations between these two characters.

The process of squareheading was found to shorten the average in-ternode length. The effect on density thus produced, however, is slight.

The phenotypic transmission of the squareheadedness of Dale Gloria is dependent on the type of the lax non-squarehead parent. In some crosses there was a large proportion of lax squarehead forms in the \(F_{2}\), while in others there were none of these forms.

Although in \(F_{2}\) progenies resulting from dense and lax crosses an almost perfect correlation exists between rachis length and density, these two characters are not necessarily correlated. Rachis length is the indirect product of average intemode length and number of internodes. The correlation between density and rachis length becomes less and less as the difference between the number of internodes of the parental forms increases.

High degrees of correlation were found between average internode
length and lensth of culn, length of sterile glames, and length of kernels, and average rachilla internode length. These, together with other obervatons, show that density and the shortening of these other length characters are the result of a single dwarfing factor.

Plants exhibiting the spelt character are not as much affected by the density factor as are those that show satizum characters.

Compact forms may be produced by crossing a lax sativum with lax forms of other species. Dense forms may also appear occasionally in crosses where neither parent is a satizum. Compactness is not a characteristic of sativum forms; other species also may exhibit this character.

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\footnotetext{
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